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Nucleic Acid Array consisting of Selective Monocyte Macrophage Genes

Description

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The present invention relates to an array consisting of oligo- or polynucleotide probes that are applied and immobilised on a solid substrate. The array is characterised in that sequences of a selection or all of the selective monocyte macrophage genes are fixed on the surface, which are enlisted in Tables 1 to 6. This nucleic-acid array permits the diagnosis of the rheumatoid arthritis, associated analysis of the efficiency of treatment and the monitoring of side effects occurring in the anti-tumour necrosis factor (TNF) therapy and hence the selection of the therapeutic approach that is most efficient for the respective patient suffering from rheumatoid arthritis. The present invention moreover relates to a nucleic-acid array for prognosis and for the development of novel pharmaceuticals with an anti-TNF orientation or of such pharmaceuticals that interfere with the control loop thereof.

The cells of the monocyte/macrophage system play a substantial role in the activation of and in maintaining inflammation cascades in the blood and in the tissue, e.g. in the course of rheumatoid arthritis, and also in other chronic inflammatory diseases as well as in auto-aggressive conditions. In the case of these diseases monocytes and macrophages are highly activated, display variations in the occupation of their surface molecules, enter into contact with other cells and secrete certain messenger substances such as those including TNF alpha, which contribute to a sustained inflammatory process. TNF alpha is a cytokine

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formed by monocytes / macrophages, lymphocytes and mastocytes, which takes an influence on inflammation, sepsis, lipid and protein metabolism, haematogenesis, angiogenesis, the healing of wounds and a defensive immune response and which produces also a cytolytic or cytostatic effect on tumour cells.

In the event of inflammatory diseases, monocytes / macrophages display a characteristic, pathologically varied gene
expression pattern with distinct variations, compared
against healthy probands. Bio-information methods known to
those skilled in the art, e.g. the significance and cluster
analysis, it is possible, inter alia, to identify or diagnose genes with a similar behaviour and highly or low-regulated genes from the hybridisation patterns of a nucleicacid array.

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With the ever-increasing availability of the high-throughput methods in the form of nucleic-acid arrays, with the exponentially growing information about the human genome and gene expression, as well as with the global networking of data bases storing structured biomedical information, the approach to and the view of chronic inflammatory and inflammatory-rheumatic conditions has been fundamentally changed. The molecular processes can be defined by application of the improved understanding of the molecular bases of cell-, tissue- and disease-specific gene expression whilst they contribute to enabling an early diagnosis and improved prognosis. On the other hand, micro-array technologies ensure the development of more efficient forms of therapy of the rheumatoid arthritis and other chronic inflammatory diseases, permitting a high-rate screening system. Moreover, these multiple techniques allow for the acWO 2004/016809 3 PCT/DE2003/001822

celerated development of pharmaceutical and biologically effective medicaments (biologicals) and also a more rapid assessment of the side effects of medicaments. For this reason, this method entails an economic profit, also in terms of national economics.

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The micro-array technology constitutes a miniaturisation of analytic processes on the basis of DNA or RNA hybridisation in a high-throughput method. At the same time, it is possible to analyse thus many thousands of different DNA/DNA (DNA/RNA) interactions within a single test assay. mRNA expression profiles are determined by means of DNA arrays by hybridisation of labelled cRNA or cDNA samples. These technologies require a high degree of automation and standardisation, with the establishment and utilisation of appropriate sample and data bases (sequence information, oligo-nucleotides). The presently employed DNA arrays are distinguished by their substrate material (nylon membranes, glass surfaces, precious metal, vapour-deposited glass surfaces, synthetic materials), by the length or by the production of the DNA sequences immobilised on the substrate and by the labelling technique applied for a sample to be fixed. By way of analogy to the methods of DNA hybridisation in the Southern/Dot-blot test, it is possible to fix DNA sequences on a filter in punctiform shape and in a systematic succession, by spotting by means of a pressure head, by piezo printing methods (ink-jet technology) or by photolithographic methods (chemical direct synthesis on the substrate material). The DNA may be a cDNA, a PCR product or a synthetically produced oligo-nucleotide. Each of these sequences so applied is hence assigned to a specific site in a known array. The RNA may be decontaminated from a clinical sample or also a sample for pharmaceutical examination

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and, after transcription by reverse transcription, may be hybridised with the complementary nucleic-acid strands present on the array, which are applied in a high genome-encompassing number or also in a pre-selected number. The sample is labelled by means of integrated radio-active nucleotides, via biotin/streptavidin interactions, via amplifications of the digoxigenin enzyme or also via direct or indirect integrated fluorescence dyes. The information is read from the intensity of radioactivity or from fluorescence on a specific site of the substrate material and hence permits the conclusion of the particular relative quantity of the specifically linked DNA or RNA sequence, which had been present in the sample so labelled.

The activation and deactivation of genes constitutes the basis of all biologic processes and furnishes moreover an extremely sensitive response to varied outside conditions. The extraction of RNA form a biologic sample, the action of labelled cDNA or RNA upon a nucleic-acid array (hybridisation) and its analysis allow for the derivation of a great number of parts of information, with a minimum of time, about the condition of the cells in the biologic sample under varied conditions. The technology based on the hybridisation of nucleic acids entails the advantage of an extremely high specificity, sensitivity and comparatively easy high-speed feasibility.

When genes are activated or deactivated in monocytes / macrophages in a non-physiologic manner, this may be the cause of inflammatory diseases or a measurable sign of such diseases. In an ideal case, the therapy with medicaments producing an anti-TNF effect should normalise the pathol-

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ogically varied gene expression in the respective cells to the level of that found in healthy patients.

From the examination of the gene expression profiles one could expect that a novel molecular characterisation of the rheumatoid arthritis and other chronic inflammatory diseases becomes possible and hence a classification by subgroups can be established by patho-physiological peculiar characteristics. With the anti-inflammatory anti-TNF therapies, one can therefore expect prognostic forecasts about the level of aggressiveness in the future development. This would take an early influence on the selection and intensity of the drug-based therapy with the medicaments so far known and used as therapeutic in the case of chronic in-15 flammations, and also with biologically effective TNF blockers. On the other hand, this leads to further starting points for scheduling the form of therapy in view of the potential side effects caused by the action of these medicaments, whilst it is possible to estimate the effects of the side effects at an early stage.

On account of therapies with an anti-TNF orientation in cases of rheumatoid arthritis and other chronic inflammatory or auto-aggressive diseases, firstly a potential genesis of neoplastic changes up to the formation of tumours is under discussion, and secondly, the anti-TNF therapy reduces the defensive immune response so that infections occur more often in treated patients, inter alia tuberculosis.

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When nucleic-acid array systems are used it is possible to check the expression of tumour-relevant genes in the course of anti-TNF treatment and hence early indications of potenWO 2004/016809 6 PCT/DE2003/001822

tial neoplastic changes can be obtained so that it becomes possible to counteract an incipient tumour formation at an early stage and the anti-TNF therapy can be appropriately adapted or, if necessary, stopped.

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The invention is based on the problem of providing means for monitoring the efficiency and also the side effects of the anti-TNF therapy and for permitting also the fine diagnosis of an inflammatory disease and hence the selection of that form of therapy, which is most efficient for the respective patient. Another problem underlying the present invention consists in the follow-up monitoring of the efficiency and the side effects of novel pharmaceuticals with anti-TNF orientation within the scope of clinical studies. In accordance with the present invention, a new array is provided that consists of oligo- or poly-nucleotide probes that are applied and immobilised on a solid substrate. Compared against genome-wide DNA chips so far known, the advantage of the present invention resists in cost savings in the production of the nucleic-acid array because it contains predominantly only genes that are of interest for a solution to the problem of the invention, which minimises

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In accordance with the present invention, the problem is solved by a nucleic-acid array including sequences of a selection or of all of the selective monocyte macrophage genes mentioned in Tables 1 to 6, which are applied on the surface of the array. With reference to the name of the gene or the sequence, respectively, or the accession number it is possible to determine the sequence from data bases accessible to the general public, preferably GeneBank or

the expenditure in terms of data analysis and evaluation

and reduces the costs thereof.

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EMBL. The sequences of the nucleic acids present on the array may consist of genes whose level of expression is changed by a therapy involving an Anti-TNF effect.

If necessary, further genes may be present on the inventive nucleic-acid array, expediently those which are known to be expressed in each cell and to constitute part of the basic genotype of the cell. The genes coding for these nucleic acids are usually referred to as house-keeping genes and are used for standardisation of the signals obtained. The array may contain the aforementioned sequences in the form of DNA, complementary RNA or chemically modified nucleic acids, preferably PNS (protein nucleic acid).

The genes or gene sequences may be selected genes of rheumatoid arthritis or other chronic inflammatory diseases, which are relevant for the disease or the side effects, selected preferably from the monocyte/macrophage cell system. If applicable, alleles, derivatives and/or splicing variants of the genes or gene fragment sequences or oligomer sequences may be present on the surface of the array. The congruence of the sequences on the array with the corresponding sequences in Tables 1 to 6 should be at least 80% in the protein-coding mRNA segments.

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The substrate onto which the nucleic acids are applied may be any substrate whatsoever which is usually employed for RNA or DNA arrays. The methods for the application and immobilisation of the nucleic acids are part of prior art and are common to those skilled in the art. For coupling to the aforementioned sequences, the substrate may be coated with reactive groups, metal compounds or alloys. The genes or gene sequences may be applied, for instance, by spotting

techniques, immobilising techniques or by techniques of insitu synthesis of oligomers or in an enantiomorphic manner in the form of RNA.

- The inventive array may be used, for instance, for measuring the monocyte/macrophage activation or the inflammatory activity in the blood or cell tissue in the case of inflammatory diseases, preferably rheumatoid arthritis. The array may be employed, for example, for the early diagnosis of the aforementioned diseases in genetically predisposed patients even before clinical symptoms become manifest. Another field of application is the fine diagnosis, preferably the subdivision of patients into sub-groups requiring each a different therapy or different medication. The array may furthermore be used to monitor the therapy, to follow up side effects, to establish a prognosis and to identify new pharmaceutical targets in the case of the aforementioned diseases.
- To this end, blood or tissue is sampled on the patients un-20 der examination, from which RNA is isolated by the known standard techniques and prepared, if necessary, as total RNA or poly A+-RNA for further use. By application of reverse transcriptase it is possible to transcribe the RNA into cDNA and to label it with a marker, e.g. a fluorescent 25 dye, a radioactive nuclide or an enzyme such as alkaline phosphatase. In addition, the RNA may be used in labelled or unlabelled condition for hybridisation of the nucleicacid array. After hybridisation of the array with the nucleic-acid samples and after subsequent washing steps, the 30 bonding of the sample to the sequences present on the array may be examined with any appropriate method. In the case of a fluorescent labelling, these methods are optical tech-

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niques whilst in the case of radioactive labelling of the samples auto radiography would be employed and in the case of enzyme labelling enzymatic analytical techniques are applied, e.g. the conversion of a colourless substrate into a multi-colour product.

An inverse analysis of total or mRNA with the sequences indicated in Tables 1 to 6, which is fixed to a solid phase, is equally possible. To this end, blood- or tissue-specific RNA molecules from up to 500 patients are fixed to the RNA 10 micro arrays. The qualitative / quantitative analysis of the transcript quantity of relevant genes is then carried out with the selected genes, gene segments or oligomers described in Tables 1 to 6. The RNA samples are spotted on coupling substrates and are composed of total RNA or mes-15 senger RNA. The RNA serves as target for the highly significantly expressed genes derived from DNA micro arrays according to Tables 1 to 6, which are used as labelled probes for hybridisation. The coupling of biotinylated RNA or messenger RNA on glass substrates (slides) coated with 20 streptavidin is proposed. After the RNA has been labelled with biotin derivatives the RNA is applied by spotting on glass or synthetic slides coated with poly-L-lysine or with streptavidin, with subsequent drying. RNA degradation is prevented in this manner. An alternative is the covalent 25 coupling of RNA by bonding to reactive substrate materials, in which case preferably UV irradiation is used for catalysing. Additionally, a multiple simultaneous labelling of different genes, gene units or oligomers with various labelling species such as radio activity, fluorescein, di-30 goxigenin and enzymatic markers is expedient.

Parallel different labelling of the probes with different fluorescent dyes is possible. As an alternative, enzymatic or even radioactive probe labelling should be mentioned. For a quantification and quality check, labelled house5 keeping genes (alpha, beta, gamma actin, GAPDH, etc.) are employed. The analysis is preferably carried out in this case in parallel and simultaneously with a maximum of 50 gene probes per test set-up.

10 In addition to the simplification of the biometric analysis by coupling RNA species to substrate materials, this system allows for a rapid diagnosis and offers a complex high-speed diagnosis, prognosis and therapy control on an individual basis for the patients. In particular in the case of pharmacologic development strategies, the system allows for a rapid performance at a high throughput.

The following examples and illustrates only serve for explanation and do not restrict the scope of the invention in any way whatsoever.

1. Isolation of monocytes

In the method employed here, the selection of selective highly pure monocytes of peripheral blood was applied in order to enable a finding (1) on the specificity of the disease, (2) on the application of the anti-TNF-alpha therapeutic as "biological", (3) in relation to a comparison against a healthy proband, and eventually also (4) for assessment of gene-diagnostic potentials relevant in relation to anti-TNF-alpha. For this test, the peripheral haemo-leucocytes from peripheral blood were enriched by Fikoll gradient density centrifugation. This fraction, which constitutes an individually different composition of

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monocytes (5 - 12%), CD4+ T-cells (85 - 92%), CD8+ T-cells (5 - 10%), NK cells (2 - 5%), basophilic and neutrophile granulocytes, was subjected to further purification steps for the extraction of specific monocyte fractions. Both, negative selections operating on removal of all other cell fractions via magnetic beads anti-body interactions, and positive selections by CD14+ labelling via magnetic beads, or even FACS cell grading techniques were applied. In both methods, monocyte cell purity levels of roughly 96% were achieved.

2. RNA Extraction

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The pure monocyte fractions were received in RNA lysis buffer whereupon the RNA was purified by means of a commercially available RNA purification kit (Qiagen). The RNA was transcribed into cDNA by application of established cDNA transcription methods by reverse transcription and then subjected to a further step of linear amplification by application of the "Eberwine protocol" for the extraction of aRNA (amplified RNA). The quantity and the quality of RNA, cDNA and aRNA were verified each by gel electrophoresis, photometric analysis and measurements carried out with the Bioanalyzer 2100 (Agilent company).

3. Affymetrix Chip Hybridisation Specific oligo-nucleotides derived directly from data base sequences were used as DNA samples for expression analyses in the system available from the Affymetrix company. These samples are hybridised on the array with targets from fluorescence-labelled, reversely transcribed samples in the form of cDNA or with linearly amplified samples in the form of aRNA.

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Hybridisation of the genome-wide Affymetrix array (U-133A) and subsequent processing are carried out by machine under standard conditions in compliance with the instructions by the producer, Affymetrix, using a special hybridisation and washing device with the specific buffers. After hybridisation, gene expression patterns are produced via the ratio of fluorescence intensity levels at a defined wavelength. Such high-throughput expression analyses permit comparisons of the expression quantities of genes in healthy and diseased persons at the same time or comparisons of the gene expression before and after drug addition for risk assessment (pharmaceutical/toxicity genomics), for fine diagnosis and for assessment of the complexity of diseases.

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4. Evaluation of Data

Here, aRNA samples from peripheral haemo-monocytes (1) of healthy blood donors, (2) of chronic active patients suffering from rheumatoid arthritis before treatment, and (3) after treatment with TNF-alpha antibodies were used. The success of the treatment was assessed by unambiguous laboratory clinical parameters and by the criteria of the internationally applicable parameter examinations (ACR criteria) to be applied on a clinical level. The objective and the purpose of this examination of groups of three were the establishment of characteristic gene expressions in the following group definitions:

A gene-regulatory specificity of a disease in active rheumatoid arthritis, without treatment, compared against the gene expression of healthy probands.

- The characterisation of a gene-regulatory specific interpretation of the anti-TNF-alpha treatment and an assessment of the treatment by comparison against the gene expression of the active disease without treatment and by comparison against the gene expression of the healthy probands.
- 3. Ensured assessment of side effects by the anti-TNF medicament "biological". Here, the specific gene expression of the patients suffering from rheumatoid arthritis, who are treated with anti-TNF-alpha, was compared against the gene expression of the same patients without treatment and the gene expression in healthy blood donors.

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The individual gene expressions were processed and measured within the genome-wide human Affymetrix array (U-133A) in the appertaining Affymetrix hybridisation/washing and reading system. The evaluation is carried out in 4 steps:

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 The determination of the significant genes detected in the expression analysis, e.g. by the "foldchange method" or by SAM (Significance Analysis of Microarrays).

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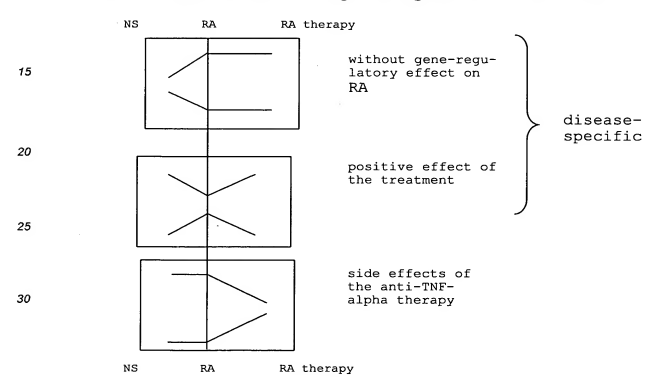
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2. The separation of the significant genes into different sub-populations on the basis of the examination of the expression characteristics of these genes by means of cluster analysis carried out by techniques such as "hierarchical clustering"!, "self-organizing maps" or "k-means clustering". 5

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- 3. The evaluation of the behaviour of the significant genes within the clusters with integration of the clinical information (rheumatoid arthritis (RA), anti-TNF therapy) and on the basis of the empirical values furnished by specialists.
- 4. The classification of the involved genes by biologic pathways.

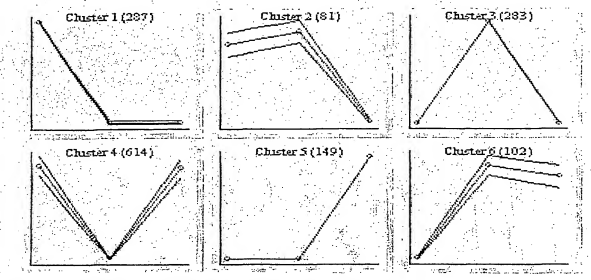
General characteristics of the significant genes within the clusters:



35 Ill. 1: Schematic representation of the cluster analysis

The gene expression behaviour of a healthy normal donor (NS) and of an active patient suffering from rheumatoid arthritis (RA) before and after anti-TNF-alpha therapy was

compared by means of a cluster analysis. The results are represented in Illustrations 1 and 2.



15 Ill. 2: Cluster analysis based on real data

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The illustrations show the gene expressions by the cluster analysis (n=6 cluster). The number of the genes involved is indicated in brackets. The result of the cluster analysis furnishes a confidence interval in addition to the mean gene expression behaviour of all the genes contained in a cluster.

The clusters displayed the following characteristics:

- 25 CLUSTER-1: Compared against a healthy proband, the disease-specific gene expression is reduced; in this case, the anti-TNF treatment does not create a generegulatory effect.
- 30 CLUSTER-2: Side effects: Represented by the effect of the medication in anti-TNF-alpha treatment, there is a reduced expression of the associated genes in the treated patient.

- CLUSTER-3: Compared against a healthy proband, the disease-specific gene expression is stronger. The anti-TNF-alpha treatment creates a positive effect.
- CLUSTER-4: Compared against a healthy proband, the disease-specific gene expression is increased. The anti-TNF treatment creates a positive effect.

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- 10 CLUSTER-5: Side effects: Represented by the effect of the medication in anti-TNF-alpha treatment, there is an increased expression of the associated genes in the treated patient.
- 15 CLUSTER-6: Compared against a healthy proband, the disease-specific gene expression is increased. The anti-TNF-alpha treatment does not create a gene-regulatory effect in this case.
- 20 Tables 1 to 6 show the genes contained in the clusters described above, together with the Affymetrix designation (left) and their defined GeneBank Accession Number, inclusive of a description.

Table 1: Genes from Cluster Analysis 1

| Affvmetrix | Dogovintion of the semience in the GeneRank data base |
|-------------|--|
| internal | מווע מכלתכווני דיון כיווי סיווים מכלת |
| designation | Translate Cds. /FER-mRNR /GEN-HIA-DOB1 / |
| 211654_x_at | gb:M17565.1 /DEE=Human MHC class 11 DX-Deta associated Williams, DEW1 Fiorein, Compress Company / Company |
| 213831_at | Consensus includes gb:X00452.1 /DEF=Human mRNA for DC classII histocompatibility antigen alpha chain: / Enrange PROD=DC classII histocompatibility antigenalpha-chain /DB_XREF=gi:32265 /UG=Hs.198253 major histocompatibility complex, |
| 212203_x_at | |
| 209480_at | gb:N16276.1 /DEF=Human HHC class II HLA-DR2-Dv12 mRNA DQv1-beta, complete cds. /FEA=mRNA /GEN=HLA-DRB2 / gb:N16276.1 /DEF=Human HHC class II HLA-DR2-Dv12 mRNA DQv1-beta, class II, DQ beta 1 /EL-gb:M60028.1 gb:M17564.1 b-No1xo 1 cb:Ng141 1 cb:N16776 1 cb:NN 002123.1 |
| 218345_at | pornation government / Inc. FEA-mRNA / Journal of the particular carcinoma-associated antigen 112 (HCA112), mRNA. / FEA-mRNA / Journal of the particular carcinoma-associated antigen 112 /DB XREF-g1:8923717 /UG-Hs.12126 hepatocellular carcinoma-associated antigen 112 /DB XREF-g1:8923717 /UG-Hs.12126 hepatocellular |
| 221491_x_at | Carcinoma-associated antigen 112 / FLEGU: AFZ-3031011 35-111 35-111 35-111 AFZ-31 A |
| | UG-Hs.279930 major nistocompatibility compres, trass 27, 5% contract for clone MGC:14507. |
| 211734_s_at | gb:BC005912.1 /DEF-Homo saplens, Fc fragment of 1gE, high affinity I, receptor 101, arpus polypeptide /DB_XREF-g1:13543505 mRNA, conpiete cds. /FEA-mRNA /PROD-Fc fragment of 1gE, high affinity I, receptorfor; alpha polypeptide /DB_XREF-g1:13543505 |
| 201330_at | EF-Homo sapiens arginyl-tRNA synthetase (RARS), mRNA. /EEA-mRNA /GEN-RARS /PROD-arginyl 8 /UG-Hs.180832 arginyl-tRNA synthetase /FL-gb:BC000528.1 gb:NM 002887.1 |
| 201114_x_at | GEN-PSNA7 / PEF-Homo sapiens proteasome (prosome, macropain) subunit, alpha type, 7 (PSNA7), mRNA. /FEA-mRNA / GEN-PSNA7 /PROD-proteasome (prosome, macropain) subunit, alphatype, 7 / DB XREF-qi:4506188 /UG-Hs.233952 proteasome GEN-PSNA7 /PROD-proteasome (prosome, macropain) subunit, alphatype, 7 / DB XREF-qi:4506188.1 gb:NM 002792.1 |
| 213418_at | (prosome, matropain) subjury of the property of the protein 6 (HSP769) (HSP769), mRNA. /FER-CDS / Consensus includes gb:NM 002155.1 /DEE-Homo sapiens heat shock 70kD protein 6 (HSP70B) / GEN-HSPA6 /PROD-heat shock 70kD protein 6 (HSP70B) / Consensus includes gb:NP protein 6 (HSP70B) / Consensus gb:NP protein 6 (HSP70B) / Consensus gb:NP prote |
| 208961_s_at | FL-going U02155.1 gb:AB017493.1 /DEF-Homo sapiens mRNA for DNA-binding zinc finger(GBF), complete cds. /FEA-mRNA / PROD-DNA-binding zinc finger(GBF) /DB XREF-gi:3582142 /UG-Hs.205313 core promoter element binding protein /FL-gb:BC000311.1 FL-DC002031 st. arginist 1 ch:BR017487.1 db:RN 001300.2 |
| 217753_s_at | gu:Browspari; gr.n.com. 3. Section |
| 201403_s_at | pb:NM 004528.1 /DEF=Homo saptens microsomal glutathione S-transferase 3 (MGST3), mRNA. /FEA-mKNA /GEA-MCA313 / PROD=microsomal glutathione S-transferase 3 /DB XREF=91:4758713 /UG=Hs.111811 microsomal glutathione S-transferase 3 / FL=ap:BC000505.1 qb:BC003034.1 gb:AF026977.1 gb:NM 004528.1 |
| 203103_s_at | gb:NM 014502.1 /DEF=Homo sapiens nuclear matrix protein NMP200 related to splicing factor PRE19 (NMP200), mixw. /GEN=NMP200 /PROD=nuclear matrix protein NMP200 related tosplicing factor PRR19 /DB_XREF=gi:7657380 /UG=Hs.173980 nuclear matrix protein NMP200 related to splicing factor PRR19 /FL=gb:NM 014502.1 |
| 221903_s_at | Consensus includes gb:BE046443 /FEA-EST /DB_XREF-gi:8363496 /DB_XREF-esE:hn47d10.xz /CLONE-IN44E:30z0003 /UG-ns.1002/ KIAA0849 protein |

| 217379_at | .121934 /DEF=Human DNA .10) pseudogene, ESTs, chromosome 6. Contain |
|-------------------|---|
| 206120_at | sapiens s.83731 C |
| 202737_s_at | sapiens U6 snRNA-associated ::6912485 /UG=Hs.76719 U6 snRN 21.1 gb:AF251218.1 |
| 201416_at | 528420 /FEA-EST /DB XREF=g1:13519957 ermining region Y)-box 4 /FL-gb:NM 0 |
| 214084_x_at | Consensus includes gb:AW072388 /FEA=EST /DB XREF=g1:6027386 /DB XREF=est:xa07d05.x1 /CLONE=IMAGE:2567625 /UG-Hs.1583 |
| 204861_s_at | sapiens baculoviral IAE repeat-containing RE-q1:4758751 /UG-Hs. 19019 baculoviral IAE |
| 221666_s_at | sapiens, clone MGC:10332, mRNA, complete cds. /FEA=mRNA /PROD=U -Hs.71869 apoptosis-associated speck-like protein containing a C |
| 218421 <u>a</u> t | sapiens hypothetical protein FLJ22239 (FLJ32329), mRNA. /FEA-mRNA 32440 /UG-Hs.34516 hypothetical protein FLJ23239 /FL-db:NN 022766.1 |
| 217794_at | |
| 201558_at | sapiens RAEI (RNA export 1, S.pombe) homolog (RAEI), mRNA. /FEA-mRNA /GEN-RAEI |
| 218055_s_at | /DEF=Homo |
| 202191_s_at | 439987 /FEA-EST /DB_XREF-g1:9439470 /DB_XREF-est:HTH1-745F /UG-Hs.226133 growth arre 05890.1 |
| 205550_s_at | gb:NM 004899.1 /DEF=Homo sapiens brain and reproductive organ-expressed (TNFRSF1A modulator) (BRE), mRNA. /FEA-mRNA / GEN-BRE /PROD-brain and reproductive organ-expressed (TNFRSF1Amodulator) /DB XREF=g1:4757871 /UG-Hs.80426 brain and reproductive organ-expressed (TNFRSF1A modulator) /EL-db:BC001251.1 db:NM 004899.1 db:L38616.1 |
| 202941_at | o sapiens NADH dehydrogenase (ubiquinone) flavoprotein 2 (24): hydrogenase (ubiquinone) flavoprotein 2 (24):D) /DB XREF=g1: n 2 (24):D) /FL=gb:NM 021074:1 gb:BC001632.1 qb:N22538.1 |
| 217814_at | sapiens GK(1s.8207 GK001 |
| 212051_at | 676803 /FEA=EST /DB XREF=91:2657325 /DE CDNA: FLJ23260 fis, clone COLO5804, hi |
| 212386_at | |
| 218571_s_at | sapiens HSPC134 s.279761 HSPC134 |
| 203462_x_at | <pre>jb:NM 003751.1 /DEF=Homo saplens eukaryotic translation initiation factor 3, subunit 9 (eta, 116kD) (EIF3S9), mRNA. / FER=mRNA /GEN=EIF3S9 /PROD=eukaryotic translation initiation factor 3, subunit 9 (eta, 116kD) /DB KREF=g1:4503526 / UG=H8:57783 eukaryotic translation initiation factor 3, subunit 9 (eta, 116kD) /FI=cb:UG>S83.1 ob:NN 003751.1</pre> |
| 218642_s_at | sapiens hypothetical protein MGC2217 (MGC2217), mRNA: /FEA-mRNA /GEN= n MGC2217 /DB_XREF=gi:13236525 /UG-Hs.323164 hypothetical protein MGC2 |
| 200024_at | gb:NM 001009.1 /DEF=Homo sapiens ribosomal protein S5 (RPS5), mRNA. /FEA-mRNA /GEN-RPS5 /PROD=ribosomal protein S5 / DB XREF=gi:1506728 /UG-Hs.76194 ribosomal protein S5 /FL-gb:NM 001009.1 gb:U14970.1 |
| 218101_s_at | gb:NM 004549.1 /DEF-Homo sapiens NADH dehydrogenase (ubiquinone) 1, subcomplex unknown, 2 (14.5kD, B14.5b) (NDUFC2), mRNA. /FER-mRNA /GEN-NDUFC2 / PROD-NADH dehydrogenase (ubiquinone) 1, subcomplex:unknown, 2 (14.5kD, B14.5b) / DB_XREF-gp::4758783 /UG-Hs.193313 NADH dehydrogenase (ubiquinone) 1, subcomplex unknown, 2 (14.5kD, B14.5b) / EL-gp:AF087659.1 gb:AF070652.1 gb:M1 004549.1 |
| | |

| | sapiens, clone MGC:2198, mRNA, complete cds. /FEA=mF Hs.227152 mannan-binding lectin serine protease 1 (C 1 |
|-------------|---|
| 208454_s_at | gb:NM 016134.1 /DEF-Womo sapiens aminopeptidase (LOC51670), mRNA. /FER-CDS /GEN-LOC51670 /PROD-aminopeptidase / DB XREF-gi:7706386 /UG-Hs.278993 aminopeptidase /FL-gb:RF107834.1 gb:NM 016134.1 |
| 214351_x_at | 189278 /FEA=EST /DB_XREF=g1:2849396 |
| 207075_at | gb:NM 004895.1 /DEE-Homo sapiens chromosome 1 open reading frame 7 (ClORF7), mRNA. /FEA-mRNA /GEN-CLORF7 / PROD-Chromosome 1 open reading frame 7 /DB_XREF-q1:4757727 /UG-Hs.159483 chromosome 1 open reading frame 7 / FL-qb:AF054176.1 gb:NM 004895.1 |
| 201449_at | Consensus includes gb:AL567227 /FEA-EST /DB XREF=g1:12920378 /DB XREF=est:AL567227 /CLONE-CSODF027YA11 (3 prime) / UG-Hs.239489 TIA1 cytotoxic granule-associated RNA-binding protein /FL-qb:NM 022037.1 gb:M77142.1 |
| 213720_s_at | Consensus includes gb:AI831675 /FEA=EST /DB_XREE=g1:5452346 /DB_XREE=est:#J50g03.x1 /CLONE=IMAGE:2406292 /UG=Hs.78202 SNISNF related, matrix associated, actin dependent regulator of chromatin, subfamily a, member 4 |
| at | Consensus includes gb:AV716798 /FEA-E5T /DB XREF=gi:10813950 /DB XREF=est:AV716798 /CLONE-DCBAKB02 /UG-Hs.6375 uncharacterized hypothalamus protein HT010 /FL-gb:AF220184.1 gb:NM 018471.1 |
| 208095_s_at | ens calciumcalmodulin-dep ilciumcalmodulin-dependent in-dependent protein kinas |
| at | gb:NM 006600.1 /DEF-Homo sapiens nuclear distribution gene C (A.nidulans) homolog (NUDC), mRNA. /FEA-mRNA /GEN-NUDC / PROD-nuclear distribution gene C (A.nidulans)homolog /DB_XREF-gl:5729952 /UG-Hs.263812 nuclear distribution gene C (A.nidulans) homolog /FL-gb:BC002399.1 gb:BC003132.1 gb:ĀB019408.1 gb:AF130736.1 gb:AF125465.1 gb:AF100760.1 gb:NM 006600.1 |
| | sapiens TRK-fused gene (TFG), mRNA /FEA-mRNA /GEN-TFG /PROD-TRK-fused gene /DE .ne /FL-gb:NM 006070.1 |
| 9t . | gb:NM 003993.1 /DEF=Homo sapiens CDC-11ke kinase 2 (CLK2), transcript variant phc1k2, mRNA. /FEA=mRNA /GEN=CLK2 / PROD=CDC-11ke kinase 2 isoform hc1k2 /DB XREF=g1:4502882 /UG=Hs.73986 CDC-11ke kinase 2 /FL=gb:NH 003993.1 gb:L29218.1 |
| | sapiens KIAA0274 gene product (KIAA0274), mRNA: /FEA-mRNA /GEN=KIAA0274 /F 133 /UG-Hs.10037 KIAA0274 gene product /FL-gb:D87464.1 gb:NM 014845.1 |
| | ojens syntaxin 18 (STX18); mRNA. /FEA-mRNA /GEN-STX18 /PROD-syntaxin 18 /DB_XREF-gi:8394375 / -gb:AB028741.1 gb:NM 016930.1; |
| | Consensus includes gb:AA382102'/FEA-EST /DB_XREF-gi:2035020 /DB_XREF-est:EST95939 /UG-Hs.3462 cytochrome c oxidase subunit VIIc |
| 201214_s_at | gb:NM 002712.1 /DEF-Homo sapiens protein phosphatase 1, regulatory subunit 7 (PPPIR7), mRNA. /FEA-mRNA /GEN-PPPIR7 / PROD-protein phosphatase 1, regulatory subunit 7 /DB_XREF=g1:4506012 /UG-Hs.36587 protein phosphatase 1, regulatory subunit 7 /FL-gb:BC000910.1 gb:NM 002712.1 |
| 216505_x_at | Consensus includes gb:AL118502 /DEF-Human DNA sequence from clone RP11-3711.9 on chromosome 20 Contains a novel gene, a gene similar to the gene for ribosomal protein 510, ESTs, STSs, GSSs and CpG islands /FEA-mRNA_3 /DB XREF-gi:8894621 / UG-Hs.284199 Human DNA sequence from clone RP11-3711.9 on chromosome 20 Contains a novel gene, a gene similar to the gene for ribosomal protein 30, ESTs, STSs, GSSs and CpG islands |
| 203459_s_at | gb:NM:022575.1 /DEE=Homo sapiens vacuolar protein sorting 16 (yeast homolog) (VPS16), mRNA. /FEA=mRNA /GEN=VPS16 / PROD=vacuolar protein sorting 16 (yeast homolog) /DB_XREF=gi:12007657 /UG=Hs.302441 vacuolar protein sorting 16 (yeast homolog) /FL=gb:AF308801.1 gb:NM 022575.1 |
| 202880_s_at | trin homology, Sec7 and colledcoil domains l(cytohesin 1/PROD=cytohesin 1, isoform 1/DB XREF=g1:4758963 /UG=Hs. 1) /FL=gb:M85169.1 gb:NM 004762.1 gb:NM 017456.1 |
| s_at | :02h12.xl |
| 203519_s_at | <pre>sapiens regulator of nonsense transcripts 2; DKF2F434D222 protei c of nonsense transcripts 2 /DB XREF-gi:11693131 /UG=Hs.3862 regu- gb:AF301013.1 gb:NM 015542.1 gb:AY013249.1 gb:AF318574.1</pre> |
| 206214_at | gb:NM 005084.1 /DEF-Homo sapiens phospholipase A2, group VII (platelet-activating factor acetylhydrolase, plasma) (FEA2G7), mRNA, /FEA-mRNA /GEN-PLA2G7 /PROD-phospholipase A2, group VII (platelet-activatingfactor acetylhydrolase, plasma) /DB XREF-gi:4826883 /UG-Hs.93304 phospholipase A2, group VII (platelet-activating factor acetylhydrolase, |

| | plasma) /FL-gb:U24577.1 gb:NM_005084.1 gb:U20157.1 |
|---------------|---|
| 209933_s_at | <pre>gb:AF020314.1 /DEF-Homo sapiens CMRF-35-H9 mRNA, complete cds. /FEA=mRNA /PROD=CMRF-35-H9 /DB_XREF=gi:4103065 / UG=Hs.9688 leukocyte membrane antigen /FI=gb:AF020314.1</pre> |
| 202105_at | ins immuno cotein 1 / |
| 218138_at | gb:NM 018848.1 /DEE-Homo sapiens McKusick-Kaufman syndrome (NKKS), mRNA. /FEA-mRNA /GEN-MKKS /PROD-McKusick-Kaufman syndrome protein /DB XREF-q1:9055271 /UG-Hs.46743 McKusick-Kaufman syndrome /FL-gb:AF275813.1 gb:AF221992.1 gb:AF221993.1 gb:NM 018848.1 |
| 220864_s_at | gb:NM 015965.1 /DEF=Homo sapiens CGI-39 protein; cell death-regulatory protein GRIM19 (LOC51079), mRNA. /FEA=mRNA / GEN=LOC51079 /PROD=CGI-39 protein; cell death-regulatory proteinGRIM19 /DB XREF=gi:7705733 /UG-Hs.279574 CGI-39 protein; cell death-regulatory protein GRIM19 /FL=qb:AF132973.1 qb:AF155662.1 qb:NM 015965.1 |
| 213355_at | Consensus includes gb:AI989567 /FEA-EST /DB_XREF-gi:5836448 /DB_XREF-est:ws34e03.x1 /CLONE-IMAGE:2499100 /UG-Hs.34578 alpha2,3-sialyltransferase |
| 212587_s_at | Consensus includes gb:AI809341 /EEA-EST /DB_XREF-gi:5395907 /DB_XREF-est:we96c09.x1 /CLONE=IMAGE:2348944 /UG-Hs.170121 protein tyrosine phosphatase, receptor type, C |
| 205992_s_at | gb:NM 000585.1 /DEF-Homo sapiens interleukin 15 (IL15), mRNA. /FEA-mRNA /GEN-IL15 /PROD-interleukin 15 / DB XREF-g1:10835152 /UG-Hs.168132 interleukin 15 /FL-gb:NM 000585.1 qb:U14407.1 |
| 203262_s_at | gb:NM 004699.1 /DEE-Homo sapiens DNA segment on chromosome X (unique) 9928 expressed sequence (DXS9928E), mRNA. / EEA-mRNA /GEM-DXS9928E /PROD=XAP-5 protein /DB XREF-qi:4758219 /UG-Hs.54277 DNA segment on chromosome X (unique) 9928 expressed sequence /FL-gb:BC000028.1 gb:DB3260-1 gb:AD001530.1 gb:NM 004699.1 |
| 213357_at | Consensus includes gb:AV701318 /FEA-EST /DB XREF-gi:10717648 /DB_XREF-est:AV701318 /CLONE=ADAAGD10 / UG-Hs.278626 ArgAbl-interacting protein ArgBP2 |
| 207628_s_at . | gb:NM 017528.1 /DEF-Homo sapiens putative methyltransferase (HASJ4442), mRNA. /FEA-mRNA /GEN-HASJ4442 / PROD-putative methyltransferase /DB XREF-gi:8923713 /UG-Hs.155020 putative methyltransferase /FL-db:NM 017528.1 |
| 201527_at | A. /FER=mRNA /GEN=ATP6S14 /PROD= |
| 202282_at | |
| 213735_s_at | EYPE 11 /FL=G0:BCUUU3/Z.1 gD:BCUU08Z9.1 gD:U735_4.1.gD:U9813Z.1.gD:AF035555.1 gD:AF069134.1 gD:NM 004493.1 Consensus includes gD:AI557312 /FEA=EST /DB_XREF=g1:4489675 /DB_XREF=est:PT2.1_16_E11.r /UG=Hs.1342 cytochrome c oxidase |
| 212472_at | SOURCE VO CONSENSUS INCLUDES 9D: BE965029 /FEA-EST / DB XREF-g1:11768976 / DB XREF-est:601658812R1 / CLONE-INAGE:3886131 /UG-HS.198793 Homo sabiens cDNA: FL722463 fis. clone HRC10126 |
| 214805_at | Consensus includes gb:U79273.1 /DEF-Human clone 23933 mRNA sequence. /FEA-mRNA /DB_XREF-gi:1710239 /UG-Hs.239483 Human clone 23933 mRNA sequence |
| 218084_x_at | gb:NM 014164.2 /DEE-Homo sapiens FXXD domain-containing ion transport regulator 5 (FXYD5), mRNA. /FEA-mRNA /GEN-FXXD5 / PROD-related to ion channel /DB_XREF-gi:11612664 /UG-Hs.294135 FXXD domain-containing ion transport regulator 5 / FL-gb:NM 014164.2 gb:AF161462.1 |
| 218204_s_at | gb:NM 024513.1 /DEE-Homo sapiens FYVE and coiled-coil domain containing 1 (FYCO1), mRNA. /FEA-mRNA /GEN-FYCO1 / PROD-FYVE and coiled-coil domain containing 1 /DB_XREF=g1:13470091 /UG-Hs.257267 FYVE and coiled-coil domain containing 1 / FI=gb:NM 024513.1 |
| 200823_x_at | gb:NM 000992.1 /DEF-Homo sapiens ribosomal protein L29 (RPL29), mRNA. /FEA-mRNA /GEN-RPL29 /PROD-ribosomal protein L29 / DB XREF-gi:4506628 /UG-Hs.183690 ribosomal protein L29 /FL-qb:U49083.1 qb:NN 000992.1 qb:U10248.1 |
| 208968_s_at | hypothetical protein, clone MGC:2478, =#8.4900 hypothetical protein /FL=qb:Al |
| 208887_at | <pre>gb:BC000733.1 /DEF=Homo sapiens, eukaryotic translation initiation factor 3, subunit 4 (delta, 44kb), clone MGC:2053, mRNA, Complete cds. /FcA=mRNA /PROD=eukaryotic translation initiation factor 3, subunit 4 (delta, 44kb) /DB XREF=gi:12653882 / USGH4s.20801 eukaryotic translation initiation factor 3, subunit 4 (delta, 44kb) /FL=gb:AF094850.1 gb:BC000733.1 gb:AF020833.1 qb:U96074.1 dp:NN 003355.1</pre> |
| 214097_at | Consensus includes gb:AW024383 /FEA-EST /DB_XREF-g1:5877913 /DB_XREF-est:wv03e06.xl /CLONE-IMAGE:2528482 /UG-Hs.1948 rlbosomal protein S21 |
| | |

| 212191_x_at | Consensus includes gb:AM574664 /FER=EST /DB_XREF=gi:7246203 /DB_XREF=est:UI-HF-BLO-abw-d-10-0-UI.s1 /CLONE=IMAGE:3057859 / UG-Ms.180842 ribosomal protein L13 |
|--------------|--|
| 217957_at | gb:NM 013242.1 /DEF-Homo sapiens similar to mouse Glt3 or D. malanogaster transcription factor IIB (AF093680), NNAN. /FFA-mRNA /GEN-AF093680 /PROD-similar to mouse Glt3 or D. malanogastertranscription factor IIB //DB.782874 /UG093680 similar to mouse Glt3 or D. malanogaster transcription factor IIB //FE-qb:BC005152.1 gb:AF093680.1 gp:NM 013242.1 |
| 201658_at | Consensus includes gb:AU151560 /FEA-EST /DB XREF=g1:11013081 /DB XREF=est:AU151560 /CLONE-NT2RP2005555 /UG-Hs.242894 ADP-ribosylation factor-like 1 /FL-gb:NM 001177.2 gb:L28997.1 |
| 218123_at | gb:NN 017835.1 /DEF=Homo sapiens chromosome 21 open reading frame 59 (C210RF59), mRNA. /FEA=mRNA /GEN=C210RF59 /PROD=hypothetical protein FLJ20467 /DB XREF=g1:8923436 /UG-Hs.5811 chromosome 21 open reading frame 59. /FL-gb:NN 021254.1 gb:BC000709.1 gb:NN 017835.1 gb:AF282851.1 |
| 205241_at | gb:NN 005138.1 /DEF=Homo saptens SCO (cytochrome oxidase deficient, yeast) homolog 2 (SCO2), nuclear gene encoding mitochondrial protein, mRNA. /FEA=mRNA /GEN=SCO2 /PROD=SCO (cytochrome oxidase deficient, yeast) homolog 2 /DB XREF=q1:4826991 /UG=Hs.278431 SCO (cytochrome oxidase deficient, yeast) homolog 2 /DB XREF=q1:4826991 /UG=Hs.278431 SCO (cytochrome oxidase deficient, yeast) homolog 2 /FI=nh.NH 005138 1 |
| 203740_at | o sapiens M-phase phosphoprotein 6 (MPHOSPH6), mRNA. /FEA-mRNA /GEN-MPHOSPH6 otein 6 /DB XREF-q1:5031918 /UG-Hs.152720 M-phase phosphoprotein 6 /EL-ab.BCN05242. |
| 221263_s_at | o sapiens hypothetical protein MGC3133 (MGC3133), mRNA. /FEA=mRNA./GEN=MGC3133 |
| 218831_s_at: | sapiens Fc OD-Fc fragme t of IgG, re |
| 201400_at | ens proteasome (prosome, macropain) oteasome (prosome, macropain) subur ome, macropain) subunit, beta type |
| 203136_at | sapiens Rab acceptor 1 (prenylated) (RABACI), mRNA5453959 /UG-Hs.11417 Rab acceptor 1 (prenylated) /F) |
| 205382_s_at | o sapiens D component of complement (adipsin) (DF), mRNA. /FEA-mRNA /GEN-DF factor D precursor /DB_XREF-gi:4503308 /UG-Hs.155597 D component of compleme 1928.1 |
| 208714_at | gb:AF092131.1 / DEFeHomo sapiens.51kDa subunit of NADH dehydrogenase mRNA, complete cds. /FEA-mRNA /PROD=51kDa subunit of NADH dehydrogenase / DB XREFeqi:5138911 /UG-Hs.7744 NADH dehydrogenase (ubiquinone) flavoprotein 1 (51kD) /FI-gb:AF053070.1 gb:AF092131.1 gb:NM 007103.1 |
| 209224_s_at | gb:BC003674.1 /DEF=Homo saplens, NADH dehydrogenase (ubiquinone) 1.alpha subcomplex, 2 (8kD, B8), clone MGC:12315, mRNA, complete cds. /FEA=mRNA /PROD=NADH dehydrogenase (ubiquinone) 1 alphasubcomplex, 2 (8kD, B8) /VG=Hs.163867 NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 2 (8kD, B8) /FEE=gi:13277539 /VG=Hs.163867 NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 2 (8kD, B8) /FEE=gi:BC003674.1 gb:AF047185.1 qb:NN 002488.1 qb:AF077029.1 |
| 210574_s_at | gb:AF241788.1 /DEF-Homo sapiens NPD011 (NPD011) mRNA, complete cds: '/FEA-mRNA /CEN-NPD011 /PROD-NPD011 / /DB XREF-gi:12005492 /UG-Hs:263812 nuclear distribution gene C (A.nidulans) homelog /EL-ob:AF241788.1 |
| 214259_s_at | 1144075 /FEA-EST /DB XREF-g1:3665884 /DB XREF-est:g163e12 xl eductase family 7, member A2 (aflatoxin aldehyde reductase) |
| 209234_at | Consensus includes gb:BF939474 /FEA-EST /DB XREF-gi:12356794 /DB XREF-est:nac76d12.x1 /CIONE-IMAGE:3440303 /UG-Hs:129908 KIAA0591 protein /FL-gb:BC001415.1 |
| 203654_s_at | gb:NM 004645.1 /DEF=Homo sapiens coilin (COIL), mRNA. /FEA=mRNA /GEN=COIL /PROD=coilin /DB_XREF=gi:4758023 /UG=Hs.966 coilin./ FL=gb:U06632.1 gb:NM 004645.1 |
| 201129_at | gb:NM-006276.2 /DEF-Homo sapiens splicing factor, arginineserine-rich 7 (35kD) (SFRS7), mRNA. /FEA-mRNA /GEN-SFRS7 / PROD-splicing factor, arginineserine-rich 7 (35kD) /DB XREF-g1:6857827 /UG-Hs.184167 splicing factor, arginineserine-rich 7 (35kD) /FL-gb:BC000997.2 gb:L22253.1 gb:NM 006276.2 |
| 202451_at | gb:BC000365.1 / DEF-Homo sapiens, general transcription factor IIH, polypeptide 1 (62kD subunit), clone MGC:8323, mRNA, Complete cds. / FEF-mRNA / PROD-general transcription factor IIH, polypeptide 1(62kD subunit). / DB_XREF-gi:12653194 / MG-Hs.89578 general transcription factor IIH, polypeptide 1 (62kD subunit) / FL-qb:BC000365.1 qb:M95809.1 qb:M95809.1 qb:MN 005316.1 |
| 218072_at | o sapiens HSPC166 protein (HSPC166), mRNB. /FEA-mRNĀ /GEN-HSPC -Hs.279836 HSPC166 protein /FL=gb:AL136688.1 gb:AF161515.1 gb: |

| 203063 at lar | |
|----------------------|--|
| | gb:NM_014634.1 /DEF=Homo sapiens KIAA0015 gene product (KIAA0015), mRNA. /FEA=mRNA /GEN=KIAA0015 /PROD=KIAA0015 gene product /DB XREE=gi:7661861 /UG=Hs.278441 KIAA0015 gene product /FL=qb:D13640.1 qb:NM 014634.1 |
| | 055366 /FEA=EST /DB XREF=g1:10809262 /DB_XREF=est:7j78f10.x1 /CLONE=IMAGE:3392587 1 differentiation, Tysophosphatidic acid G-protein-coupled receptor, 2 11.1 gb:RM 001401.1 |
| 6 | gb:NM_024952.1 /DEF-Homo sapiens hypothetical protein FLJ20950 (FLJ20950), mRNA. /FEA-mRNA /GEN-FLJ20950 /PROD-hypothetical protein FLJ20950 /DB XREF-q1:13376436 /UG-Hs.285673 hypothetical protein FLJ20950 /FF-mh:NM 024952 1 |
| 204158_s_at gb | gb:NM 006019.1 /DEF=Homo sapiens T-cell, immune regulator 1 (TCIRG1), mRNA. /FEA=mRNA /GEN=TCIRG1 / PROD=ATPase, H+ transporting, 116kD /DB XREF=qi:5174620 /UG-Hs.46465 T-cell, immune regulator 1 /FL=ob:U45285.1 ob:NM /Off019 1 |
| at | 3787 /FEA=EST /DB_XREF= L |
| at. | sapiens is.16134. g |
| | aplens cDNA FLJ13972 fis, clone Y79AA1001548, IASE ALPHA (EC 2.7.1.67). /FEA-mRNA /DB_XREF-gi catalytic, alpha polyneptide |
| 209858_x_at gt PP | |
| | nan N-sulphoglucosamine |
| 201871_s_at gb | gb:NM 015853.1 /DEF-Homo sapiens ORF (LOC51035), mRNA. /FEA-mRNA /GEN=LOC51035 /PROD=unknown protein LOC51035 / DB XREF-g1:1705653 /UG-Hs.77868 ORF /FL-qb:EC000902.1 qb:M68864.1 db:NM 015853.1 |
| | 149944.1 /DEF-Homo sapiens mRNA; cDNA DKF? 322 /PROD-hypothetical protein /DB XREF-cj |
| 217802_s_at 9b PB | iens similar to rat nuclear ubiquitous casein kinase 2 (NUCKS), mRNA. /FE ubiquitous casein kinase 2 /DB XREF-gi:12232386 nuclear ubiquitous casein kinase 2 /FF-m-h:NM 022731 1 AH-R-GOORGS 1 |
| _at |) sapiens KIAA0916 protein (KIAA0916); mRNA. /FEA-mRNA /GEN- 1s.151411 KIAA0916 protein /FL-qb:AF075587.1 qb:AF083244.1 |
| | IMAGE-2209217 / |
| at | gb:AF067854.1 /DEF-Homo sapiens adenylosuccinate lyase (ADSL) mRNA, alternatively spliced, complete cds. /FEA-mRNA 'GEN-ADSL /PROD-adenylosuccinate lyase /DB XREF-qi:3211983 /UG-Hs 7557 adenylosuccinate lyase /Fr-nh.bfn67854) |
| _at | sapiens ubiquitously-expressed transcript (UXT), mRNA. /FEA-mRNA /GEN-UXT. ssed transcript /DB XREE-gi:4759297 /UG-Hs.172791 ubiquitously-expressed transcript 092737.1 gb:NM 004182.1 gb:AF083241.1 gb:AF083242.1 |
| | saplens glutathione peroxidase 4 (ph lase 4 /DB_XREF=gi:4504106 /UG=Hs.2706 |
| at . | gb:NM 003611.1 /DEE-Homo sapiens chromosome X open reading frame 5 (CXORFS), mRNA. /FEA-mRNA /GEN-CXORFS / PAP-NA ONSET 1 PROD-Chromosome X open reading frame 5 /DB XREE-q1:4503178 /UG-H8:6483 oral-factal-digital syndrome 1 gane /FI-sh-NA ONSET 1 |
| | L523904 /FEA=EST /DB XREF=g1:12787397 /DB XREF=est:AL523904 /CLONE-CSODC003YB07 (3 prime) / (prosome, macropain) subunit, alpha type, 2 /FL-cb:NM 002787 1 |
| | Sapiens prefoldin 2 (PFDN2), mRNA. /FEA-m /FL-gb:NM 012394.2 gb:AF165883.1 gb:AF117 |
| | 502377 /FEA-EST /DB_XREF-gi:9704785 /DB_XREF-es in 2 /FL-gb:AB000462.1 gb:AF000936.1 |
| | sapie nzyme 306.1 |
| 208776_at Co | Consensus includes gb:BF432873 /FEA-EST /DB_XREF-gi:11445036 /DB_XREF-est:7n28a02.x1 /CLONE-IMAGE:3565730 /UG-Hs.90744 proteasome |
| | Consensus includes gb;BE502377 /FEA=EST /DB XREF=gi SH3-domain binding protein 2 /FL-gb;AB000462.1 gb:NB gb:NM 005327.1 /DEF=Homo sapiens L-3-hydroxyacyl-Co PROD=L-3-hydroxyacyl-Coenzyme A dehydrogenase, shox short chain /FL-gb:BC000306.1 gb:NN 005327.1 Consensus includes gb:BF432873 /FEA=EST /DB_XREF=gi |

| | (prosome, macropain) 26S subunit, non-ATPase, 11 /FL-gb:BC000437.1 gb:BC004430.1 gb:AB003102.1 gb:AF001212.1 gb:NA_002815.1 |
|-------------|--|
| 214765_s_at | KO24677.1 /DEF=Homo sapiens cDNA 437016 /UG=Hs.264330 N-acylsphin |
| 204225_at | is histone deacetylase 4 (HDAC4), mRNA. /FEA=mRNA /GEN=HDAC4 /PROD=histone O histone deacetylase 4 /FL=gb:NM 006037.2 qb:AF132607.1 |
| 214170_x_at | A669797 /FEA=EST /DB_XREF=qi:2631296 /DB_XREF=est:ag36c01.s1 /CLONE=IMAGE:1118880 |
| 218210_at | gb:NN 024619.1 /DEF-Homo sapiens hypothetical protein FLJ12171 (FLJ12171), mRNA. /FEA-mRNA /GEN-FLJ12171 /PROD-hypothetical protein FLJ12171 /DB XREF-gi:13375839 /UG-Ks.31431.hypothetical protein FLJ12171 /FL-gb:AL136631.1 gb:NN 024619.1 |
| 218652_s_at | gb:NM 017733.1 /DEF-Homo sapiens hypothetical protein FLJ20265 (FLJ20265), mRNA. /FEA-mRNA /GEN-FLJ20265 /PROD-hypothetical protein FLJ20265 /DB XREF-gi:8923239 /UG-Hs.7099 hypothetical protein FLJ20265 /FL-gb:BC001249.1 gb:BC000937.2 gb:NN 017733.1 |
| 201270_x_at | |
| 203906_at | *IMAGE:2307157 /UG-Hs.4764 KIA |
| 221516_s_at | hypothetical protein, clone MGC:1067, mRNA, complete cds. /FEA-mRNP) hypothetical protein /FLAgb:BC002587.1 |
| 209492_x_at | gb:BC003679.1 /DEF-Homo sapiens, ATP synthase, H+ transporting, mitochondrial FO complex, subunit e, clone MGC:12532, mRNA, complex detecteds. /FEA-mRNA FRODE-MFS-synthases, H+ transporting, mitochondrial FOcomplex, subunit e /DB_XREF-gi:13277543 / write accarage are superhase. H+ transporting mitochondrial FO complex, subunit e /FEA-B-ROOTSF9.1 |
| 212048_s_at | W24540 |
| 204214_s_at | o sapiens RAB32, member RAS oncogene family (RAB32), mRNA. F-gi:5803132 /UG-Hs.32217 RAB32, member RAS oncogene famil |
| 217796_s_at | o sapi EF=gi: |
| 203327_at | 122903 /FEA=EST /DB_XREF=gi:1137053 /DB_XREF=est:yx66e04.sl / gb:NM 004969.1 |
| 208659_at | ride channel ABP M 001288.2 |
| 222218_s_at | 4J400843.1 /DEF-Homo sapiens partial mRNA % A-mRNA /GEN-FDF03-M14 /PROD-cell surface re ceptor alpha |
| 211862_x_at | sapiens Usurpin-beta mRNA, complete cds. /FEA-CDS /PROD-Usurpin-beta regulator /FL-gb:AF015451.1 |
| 209080 x at | • sapiens PKCq-interacting protein PICOT (PICOT) mRNA, complete cds. OT /DB XREF=q1:6840952 /UG=Hs:4264*thioredoxin-like /FL=gb:BC00528 |
| 200789_at | gb:NM 001398.1./DEF-Homo sapiens enoyl Coenzyme A hydratase 1, peroxisomal (ECH1), mRNA. /FEA-mRNA /GEN-ECH1 /PROD-peroxisomal enoyl-coenzyme A hydratase-likeprotein /DB_XREF-gi:4503446 /UG-Hs.196176 enoyl Coenzyme A hydratase 1, peroxisomal /FEA-mRNA 001398 1 sh:UT16660 1 |
| 90610_at | |
| 201704_at | gb:NM 001247.1 /DEF=Homo sapiens ectonucleoside triphosphate diphosphohydrolase 6 (putative function) (ENTFD6), mRNA. / FER=mRNA /GEN=ENTPD6 /PROD=ectonucleoside triphosphate diphosphohydrolase 6(putative function) /DB_XREF=gi:4557422 / UG=Hs.12330 ectonucleoside triphosphate diphosphohydrolase 6 (putative function) /FL=gb:AF039916.1 gb:NM 001247.1 |
| 218026_at | |
| 214771_x_at | .1 /DEF-Homo sapiens cDNA: FLJ21951 fis, clone HEP04968. /FEA-mRNA /DB_XREF-gi:1043 |
| 208760_at | Consensus includes gb:ALO31714 /DEF-Human DNA sequence from clone IA16-358B7 on chromosome 16 Contains the UBE21 gene for ubiquitin-conjugating enzyme E21 (homologous to yeast UBC9), and an RPS20 (40S Ribosomal protein S20) pseudogene. Contains E3Ts, STSs. GSSs and a putative CpG is /FEA-mRNA /DB XREF-gi:4775608 /UG-Hs.84285 ubiquitin-conjugating enzyme E21 (homologous to |

| | yeast UBC9) /FL-gb:U45328.1 gb:U31933.1 gb:BC000427.1 gb:BC004429.1 gb:U31882.1 gb:U66818.1 gb:U66867.1 gb:U38785.1 gb:NM_003345.1 gb:U29092.1 |
|-------------|--|
| 212607_at | Consensus includes gb:N32526 /FEA=EST /DB_XREF=g1:1152925 /DB_XREF=est:yy11f04.s1 /CLONE=IMAGE:270943 /UG=Hs.300642 serologically defined colon cancer antigen B |
| 64418_at | Cluster Incl. A1472320:tj87c02.x1 Homo sapiens cDNA, 3 end /clone=IMAGE-2148482 /clone_end=3 /gb=A1472320 /gi=4334410 / ug=Hs.48504 /len=548 |
| 205819_at | gb:NM 006770.1 /DEE-Homo sapiens macrophage receptor with collagenous structure (MARCO), mRNA. /FEA-mRNA /GEN-MARCO / PROD-macrophage receptor with collagenous structure /DB_XREF-gi:5803079 /UG-Hs.67726 macrophage receptor with collagenous structure /FL-qb:AF035819.1 qb:NM 006770.1 |
| 218019_s_at | 9b:NM 021941.1 /DEE-Homo sapiens hypothetical protein FLJ21324 (FLJ21324), mRNA. /FEA-mRNA /GEN-FLJ21324 /PROD-hypothetical protein FLJ21324 /DB XREF-01:11345479 /UG-Hs.4746 hypothetical protein FLJ21324 /DB XREF-01:11345479 /UG-Hs.4746 hypothetical protein FLJ21324 /FIA-N-NM 021641 |
| 219220_x_at | sapiens GK002 protein (GK002), mRNA. FEA-mRNA /GEN-GK00 in, gibt protein; chromosome 3 open reading frame 5 /Fi-me |
| 212355_at | 075450 /FER-EST /DB_XREF=g1:3399805 /DB_XREF=est:oz82g10.x1 /CLONE=INAGE:1681890 / |
| 214735_at | Consensus includes gb:AW166711 /FEA=EST /DB_XREF=gi:6398236 /DB_XREF=est:xg27h02.x1 /CLONE=IMAGE:2628819 / UG-Hs:185140 KIAA0403 protein |
| 219041_s_at | gb:NM 014374.1 /DEE-Homo sapiens zinc finger protein (AP41, mRNA, /FEA=mRNA /GEN=AP4 /PROD=zinc finger protein / DB XREE=g1:7656889 /UG-Hs.90693 zinc finger protein /FL=qb:BC000363.1 ob:NM 014374.1 |
| 209110_s_at | gb:AL050259.1 /DEF-Homo sapiens mRNA; cDNA DKF25564D0782 (from clone DKF2p564D0782); complete cds. /FEA-mRNA /GEN-DKF2p564D0782 / PROD-hypothetical protein /DB_XREF-g1:4886476 /UG-Hs.170160 RAB2, member RAS oncogene family-like /FL-gb:D85757.1 gb:NA_004761.1 gb:AL050259.1 |
| 215691_x_at | Consensus includes gb:AV702994 /FEA=EST /DB_XREF=gi:10719324 /DB_XREF=est:AV702994 /CLONE=ADBAPB06 /UG=Hs.46967 HSPCO34 protein |
| 207614_s_at | gb:NM_003592.1 /DEE-Homo sapiens cullin 1 (CUL1), mRNA. /FEA-mRNA /GEN=CUL1 /PROD-cullin 1 /DB_XREF=g1:4503160 / UG-Hs.14541 cullin 1 /FL-gb:U58087.1 gb:NM 003592.1 |
| 213405_at | Consensus includes gb:N95443 /FEA-EST /DB XREF-gi:1267753 /DB_XREF-est:zbB1c12.sl /CLONE-IMAGE:310006 /UG-Hs.19180 Homo sapiens mRNA; cDNA DKF2p564E122 (from clone DKF2p564E122) |
| 208835_s_at | Consensus includes gb:AM089673 /FEA-EST /DB_XREF=gi:6047017 /DB_XREF=est:xd2lh11.x1 /CLONE=IMAGE:2594469 /UG=Hs.3688 cisplatin resistance-associated overexpressed protein /FL=qb:AB034205;1 |
| 202225_at | Consensus includes gb:RM612311 /FEA=EST /DB XREF=gi:7317497 /DB XREF=est:hg95e07.x1 /CLONE=IMAGE:2953380 /UG=Hs.306088 v-crk avian sarcoma virus C110 oncogene homolog /FL=qb:D10656.1 qb:NM 016823.1 |
| 218358_at | 95:NM 024324.1 /DEE-Homo sapiens hypothetical protein MGC11256 (MGC11256), mRMA. /FEA-mRNA /GEN-MGC11256 /PROD-hypothetical protein MGC11256 /DB XREF-gi:13236568 /UG-Hs.28029 hypothetical protein MGC11256 /FL-ab:RC002894.1 ob.NM 024324) |
| 213511_s_at | 167164 /FEA=EST /DB_XREF=g1:3700334 /DB_XREF=est:0008c10.x1 /CLONE= |
| 218133_s_at | gb:NM 021824.1 /DEF=Homo sapiens NIF3 (Ngg1 Interacting factor 3, S.pombe homolog)-like 1 (NIF3L1), mRNA. /FEA=mRNA /GEN=NIF3L1 / PROD=NIF3 (Ngg1 interacting factor 3, S.pombehomolog)-like 1 /DB.XREF=gi:11141898 /UG=Hs.21943 NIF3 (Ngg1 interacting factor 3, S.pombe homolog)-like 1 /FL=qb:RF182416.1 qb:NM 021824.1 db:AFD67513.1 db:xAFD67644.1 |
| 212007_at | Consensus includes qb:AI927512 /FEA-EST /DB_XREF-q1:5663476 /DB_XREF-est;wo90g11.x1 /CLONE-IMAGE:2462660 /UG-Hs.77495 UBX domain-containing 1 |
| 204690_at | gb:NM 004853.1 /DEF-Homo sapiens syntaxin 8 (STX8), mRNA. /FEA-mRNA /GEN-STX8 /PROD-syntaxin 8 /DB_XREF-gi:4159187 /UG-Hs.119525 syntaxin 8 /EL-gb:AF062077.1 qb:AF036715.1 qb:AF115323.1 db:NM 004853.1 |
| 210042_s_at | gb:AF073890.1 /DEE-Homo sapiens cathepsin X precursor, mRVA, complete cds. /FEA-mRNA /PROD=cathepsin X precursor / DB XREF=g1:3650497 /UG-Hs.252549 cathepsin Z /FEL=qb:AF032906.1 db:AF073890.1 db:RN 001336.1 db:AF13671 |
| 32091_at | omo sapiens mRNA for KIAA0446 protein, complete cds /cds=(3531,4286) |
| 201135_at | gb:NM 004092.2 /DEF=Homo sapiens encyl Coenzyme A hydratase, short chain, 1, mitochondrial (ECHSI), nuclear gene encoding mitochondrial protein, mRNA. /FEA=mRNA /GEN=ECHSI /PROD=mitochondrial short-chain encyl-coenzyme Ahydratase 1 precursor / DB XREF=gi:12707569 /UG=Hs.76394 encyl Coenzyme A hydratase, short chain, 1, mitochondrial /FI=ch:NM 004092 2 nh.D130A 1 |
| 220526_s_at | 224 (FLJ10024), mRNA. /FEA-mRNA /GEN-FLJ10024 /PROF |
| | |

| 202531_at | |
|--------------------|--|
| 213465_s_at | Consensus includes gb: BF718769 /FEA-EST /DB XREF-gi:12019514 /DB XREF-est:KEST72 /CLONE-S90413.NIH-107-R.abl /UG-Hs.36587 protein phosphatase 1, regulatory subunit 7 |
| 202025 <u>z</u> at | <pre>gb:NN_001607.2 /DEF=Homo sapiens acetyl-Coenzyme A acyltransferase 1 (peroxisomal 3-oxoacyl-Coenzyme A thiolase) (ACAAl), nuclear gene encoding mitochondrial protein, mRNA. /EEN=mRNA/GEN=ACAAl /PROD=acetyl-Coenzyme A acyltransferase 1 / DEXREFE=g1:659816 /UG=Hz. 1166160 acetyl-Coenzyme A acyltransferase 1 (peroxisomal 3-oxoacyl-Coenzyme A thiolase) / El=ch:RCNO0638, 1 ob:NM 001607.2</pre> |
| 203944_x_at | |
| 214290_s_at | 13324 /FEA-EST /DB_XREF-gi:4018929 /DB_XREF-est:ta77f02.x2 /CLONE-IMAGE:2050107 /UG- |
| 208415_x_at | DEF-Hom member |
| 216396_s_at | Consensus includes gb: AF131850.1 /DEF-Homo sapiens clone 24988 mRNA sequence. /FEA-mRNA /DB_XREF-g1:4406694 /UG-Hs.286027 etoposide-induced mRNA |
| 203033_x_at | sapiens atase /F |
| 203971_at | gb:NM 001859.1 /DEF=Homo sapiens solute carrier family 31 (copper transporters), member 1 (SLC31A1), mRNA. /FEA=mRNA / GEN=SLC31A1 /PROD=solute carrier family 31 (copper transporters), member 1 /DB_XREF=qi:4507014 /UG=Hs.73614 solute carrier family 31 (copper transporters), member 1 /FL=gb:UB3460.1 gb:NM 001859.1 |
| 203241_at | o sapiens UV radiation resistance as: ene /DB XREF=q1:4507860 /UG=Hs.13137 |
| 201142_at | %577698 /FEA=EST /DB XREF=g1:2355882 /DB XREF=est:nn22h05.s1 /CLONE=IAAGE:1084665 /UG-Hs.151777 factor 2, subunit 1 [alpha, 35kD] /FL=g5:BC002513.1 gb:J02645.1 gb:NM 004094.1 |
| 208818_s_at | gb:BC000419.1 /DEF=Homo sapiens, catechol-O-methyltransferase, clone MGC:8663, mRNA, complete cds. /FEA=mRNA / PROD=catechol-O-methyltransferase /DB XREF=gi:12653300 /UG=Hs.240013 catechol-O-methyltransferase /FL=gb:BC000419.1 gb:M58525.1 gb:M65212.1 gb:NM 007310.1 gb:NM 000754.2 |
| 203436_at | gb:NM 006413.1 /DEF=Homo sapiens ribonuclease P (30kD) (RPP30), mRNA. /FEA=mRNA./GEN-RPP30 /PROD=ribonuclease P (30kD) / DB XREF=g1:5454023 /UG-Hs.139120 ribonuclease P (30kD) /FL=gb:U77665.1 gb:NM 006413.1 |
| 201380_at | gb:NM 006371.1 /DEF=Homo sapiens cartilage associated protein (CRTAP), mRNA. /FEA-mRNA /GEN-CRTAP /PROD-cartilage associated protein /DE XREF=gi:5453600 /UG-Hs.155481 cartilage associated protein /FL-gb:NM 006371.1 |
| 203614_at | gb:NM 021645.1 /DEE=Homo sapiens KIAA0266 gene product (KIAA0266), mRNA. /FEA-mRNA /GEN=KIAA0266 /PROD=KIAA0266 gene product / DB XREF-gi:11063982 /UG-Hs.127376 KIAA0266 gene product /FI-gb:NM 021645.1 gb:D87455.1 |
| 208649_s_at | gb:AF100752.1 /DEF-Womo sapiens transitional endoplasmic reticulum ATPase mRNA, complete cds. /FEA=mRNA /PROD=transitional endoplasmic reticulum ATPase /DB XREF-gi:5410289 /UG-Hs.106357 valosin-containing protein /FL-gb:AF100752.1 gb:NN 007126.2 |
| 209012_at | Consensus includes gb:AV718192 /FEA=EST /DB XREF=g1:10815344 /DB_XREF=est:AV718192 /CLONE=FHTAABE08 /UG=Hs.171957 triple functional domain (PTPRF interacting) /FL=gb:AF091395.1 |
| 212100_s_at | Consensus includes gb: 293241 /DEF-Human DNA sequence from clone 222E13 on chromosome 22. Contains three novel genes, an ATP Synthase G Chain, Mitochondrial (EC 3.6.1.34) pseudogene and the DIAl gene for diaphorase (NADH) (cytochrome b-5 reductase) (EC 1.6.2.2). Contains ESTs, SISs, /FEA-mRNA 5 /DB NREF-qi:4826450 /UG-Hs.278314 hypothetical protein |
| 208756_at | eceptor interacting protein 1 mRNA, c -Hs.192023 eukaryotic translation ini 9067.1 gb:NM 003757.1 gb:AF116697.1 |
| 213034_at | .1 /DEF=Homo : .gi:4589641 /U |
| 200684_s_at | /FEA=EST 2L 3 /FL=g |
| 213019_at | |
| 218617_at | gb:NM_017646.1 /DEF=Homo sapiens tRNA isopentenylpyrophosphate transferase (IPT), mRNA. /FEA=mRNA /GEN=IPT / PROD=tRNA isopentenylpyrophosphate transferase /DB XREF=gi:8923064 /UG=Hs.288036 tRNA isopentenylpyrophosphate transferase / |

| | EL-qb:NM_017646.1 |
|-------------------|--|
| 208308_s_at | 1. " |
| 209429_x_at | <pre>gb:AF112207.1 /DEF=Homo sapiens translation initiation factor eIF-2b delta subunit mRNA, complete cds. /FEA=mRNA / PROD=translation initiation factor eIF-2b deltasubunit /DB_XREF=q1:6563201 /UG=Hs.169474 DKF2P586J0119 protein / FL=qb:BC001870.1 gb:AF112207.1</pre> |
| 221501_x_at | sapiens KIAA0220 s.251928 nuclear |
| 209161 <u>a</u> t | 184802 /FEA-EST /DB XREF=gi:3735440 /DB XREF=est:9 F016369.1 gb:U82756.1 gb:NM 004697.1 |
| 201414_s_at | 9b:NM 005969.1 /DEF=Homo sapiens nucleosome assembly protein 1-like 4 (NAPIL4), mRNA. /FEA=mRNA /GEN=NAPIL4 / PROD=nucleosome assembly protein 1-like 4 /DB_XREF=91:5174612 /UG=Hs.78103 nucleosome assembly protein 1-like 4 / FL=cb:U77456.1 ob:NM 005969.1 |
| 212361_s_at | Consensus includes qb:AW190070 /FEA=EST /DB XREF=qi:6464550 /DB XREF=est:xlllfll.xl /CLONE=IMAGE:2675949 /UG-Hs.1526 ATPase, Ca++ transporting, cardiac muscle, slow twitch 2 |
| 204084_s_at | 11687 /FEA-EST uronal 5 /FL-qb |
| 203200_s_at | gb:NM 024010.1 /DEE=Homo sapiens 5-methyltetrahydrofolate-homocysteine methyltransferase reductase (MTRR), transcript variant 2, mRNA. /FEA-mRNA /GEN-MTRR /PROD=methionine synthase reductase, isoform 2 / DE XREF-gi:13325067/ UG-Hs.153792 5-methyltetrahydrofolate-homocysteine methyltransferase reductase /FI-ch:NM 024010 1 ch:3F191014 |
| 201405_s_at | gb:NM 006833.1 /DEF=Homo saplens COP9 subunit 6 (MOV34 homolog, 34 kD) (MOV34-34KD), mRNA. /FEA-mRNA /GEN-MOV34-34KD / PROD-COP9 subunit 6 (MOV34 homolog, 34 kD) /DB_XREF=g1:5803095 /UG-Hs.15591 COP9 subunit 6 (MOV34 homolog, 34 kD) / [FL-gb:BC002520.1 gb:U70735.1 gb:NM 006833.1 |
| 218556_at | l to |
| 217756 x_at | sapiens small EDRK-rich factor 2 (SERF2), s. 323806 small EDRK-rich factor 2 /FL-qb: |
| 209206_at | |
| 212165_at | F070537.1 /DEF=Homo sapiens clone 2460 A sequence |
| 208983_s_at | |
| 202042_at | sapiens histidyl-tRNA syntheta: .77798 histidyl-tRNA synthetase |
| 202433_at | gb:NM 005827.1 /DEF=Homo sapiens UDP-galactose transporter related (UGTREL1), mRNA. /FEA=mRNA /GEN=UGTREL1 / PROD-UDP-galactose transporter related /DE_XREF=gi:5032212 /UG=Hs.154073 UDP-galactose transporter related /FL=gb:D87989.1 gb:NM 005827.1 |
| 201145_at | pb:NM 006118.2 /DEF=Homo sapiens HS1 binding protein (HAXI), mRNA. /FEA=mRNA /GEN=HAXI /PROD=HS1 binding protein / DB XREF=q1:13435355 /UG=Hs.15318 HS1 binding protein /FL=qb:NM 006118.2 qb:BC005240.1 qb:U68566.1 |
| 311975_at | 99671 /FEA=EST /DB_XREF=gi:9183419 /DB_XREF=es |
| 200918_s_at | gb:NM 003139.1 /DEF=Homo sapiens signal recognition particle receptor (docking protein) (SRER), mRNA. /FEA-mRNA /GEN=SRPR / PROD=Signal recognition particle receptor (dockingprotein) /DB_XREF=gi:4507222 /UG-Hs.75730 signal recognition particle receptor (docking protein) /EL=gb:BC001162.1 gb:NM 003139.1 |
| 203497_at | gb:NM 004774.1 /DEF-Homo sapiens PPAR binding protein (PPARBP), mRNA. /FEA-mRNA /GEN-PPARBP /PROD-thyroid hormone receptor interactor 2 /DB XREF-gi:4759265 /UG-Hs.15589 PPAR binding protein /FL-qb:AF055994.1 qb:NM 004774.1 qb:AF283812.1 |
| 209153_s_at | |
| 220966_x_at | gb:NM 030978.1 /DEF=Homo sapiens hypothetical protein similar to actin related protein 23 complex, subunit 5 (MGC3038), mRNA. / EEA-mRNA /GEN-MGC3038 /PROD-hypothetical protein similar to actin relatedprotein 23 complex, subunit 5 /DB_XREF-gi:13569955 / FL-gb:NM 030978.1 |
| | |

| 212785_s_at | 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 |
|-------------|--|
| | Consensus includes gu: Maiouioi / Emanesi / De Andrey: 1734/3/ / De Andrews: 21 / Casada - Maracissis / Casada |
| 219176_at | |
| 218679_s_at | PSZ8 protein (LOC51160), mRNA. /FEA-mRNA /GEN-LOC51160 /FROD-VPS28 protein / FSZ8 protein /FI-gb:AF316887.1 gb:AF182844.1 gb:NM 016208.1 |
| at. | gb:NM 000579.1 /DEF-Homo sapiens chemokine (C-C motif) receptor 5 (CCR5), mRNA. /FEA-mRNA /GEN-CCR5 /FROD-Chemokine (C-C motif) receptor 5 /FD SEF-91:4502638 /UG-Hs.54443 chemokine (C-C motif) receptor 5 /FD SEF-91:450964.1 gb:NM 000579.1 |
| 209770_at | gb:U90552.1 /DEF=Human butyrophilin (BTE5) mRNA, complete cds. /FEA=mRNA /GEN=BTF5 /FNOD=butyrophilin /DB_XREF=g1:2062/U3 / UG-Hs.284283 butyrophilin, subfamily 3, member Al /FL=gb:U90552.1 |
| 214268_s_at | ST /DB_XREF=gi:5421564 /DB_XREF=est:DKF2p434H1920_s1 /CLONE-DK |
| 201885_s_at | GD:NM 000399.3 /DEE-Homo sapiens diaphorase (NADH) (cytochrome b-5 reductase) (DIA1), nuclear gene encoding mitochondrial protein, transcript variant M, mRNA. (FEA-mRNA, (SEA-MRNA, (SEA-MRNA, ORGANA) (PRODE CONTROLLES) membrane-bound isoform / |
| 209688_s_at | DB XNEE-g116352226 / UGETS, 2 14164 dispinitative (NADD) 1 1/2 CONTROLL STATES (NATIONAL OF STATES) / CONTROLL STATES (NATIONAL OF STA |
| 203487_s_at | DB XKEE-91:134 / 12.4 / OG-NS.:0116 NOMO Septems Crope 24:00-many Sequence 7.2 90.50005070:1. \$\text{pi.Mi-01:5396.1 } \text{DEE-NHOM-2 sapiens Date: DEED-NHOM-3 } \text{DEE-NHOM-3 } \text{CEA-mRNA-7 GEN-DKF2P434A043 } \text{DEE-NHOM-3 } \text{DEEN-3 } \text{DEE-NHOM-3 } |
| 202155_s_at | |
| 221786_at | Consensus includes gb:BF197222 /FEA-EST /DB_XREF-gi:11085906 /DB_XREF-est:7m88b07.x1 /CLONE-IMAGE:3561949 /UG-Hs.12342 Homo sapiens Clone 24538 mRNA sequence |
| 212846_at | 15 |
| 209444_at | gb:BC001851.1 /DEF-Homo sapiens, Similar to RAPI, GTP-GDP dissociation stimulator 1, clone MGC:4525, mRNA, complete cds. / |
| | stimulator 1 /FL=gb:NM 021159.1 gb:BC001851.1 gb:BC001816.1 gb:AF215923.1 gb:AF237413.1 |
| 211937_at | Consensus includes gb:NM 001417.1 /DEF-Homo saplens eukaryotic translation intriation factor 4B (Lir4b), mkW. /Frw-Cos./ GEN-EIF4B /PROD-eukaryotic translation initiation factor 4B /DB_XREF-g1:4503532 /UG-Hs.93379 eukaryotic translation initiation factor 4B /FL-gb:NM 001417.1 |
| 201800_s_at | saplens oxysterol-binding protein 1 /DB_XREF=g1:1044 |
| 211730_s_at | sapiens, polymerase (RNA) II (DNA directed) polypeptide L (7.6kD), clone MGC: se (RNA) II (DNA directed) polypeptide L(7.6kD) /DB XREF=gi:13543491 /FL-gb:BC |
| 65635_at | sl Homo sapiens cDNA, 3 end /clone-bKF2p434M1928 /clone_end=3 |
| Į, | GD:NN 005701.1 /DEF-Homo sapiens RNA, U transporter 1 (RNUT1), mRNA. /FEA-mRNA /GEN-RNUT1 /PROD-RNA, U transporter 1 / OB XREF-gi:5031832 /UG-Hs.21577 RNA, U transporter 1 /FL-gb:AE039029.1 gb:NM 005701.1 |
| | EST /DB XREF=g1:12097014 /DB_XREF=est:602252342F1 /CLONE=IMAGE:434 e NT2RPI000851 |
| 203912_s_at | apiens deoxyribonuclease I-like 1 (DNASELLI), mRNA. /FEA-mRNA /GEN-DNASELLI /PROD-deoxyribonucl 06 /UG-Hs.77091 deoxyribonuclease I-like 1 /FL~gb:UO6846.1 gb:NM 006730.1 |
| 216996_s_at | Consensus includes gb:AK021557.1 /DEE=Homo sapiens cDNA FLJ1495 fis, clone HEMBA1001950, highly similar to Homo sapiens mkNA for KIAA0971 protein. /FEA=mRNA /DB XREE=gi:10432760 /UG=Hs.84429 KIAA0971 protein |
| 202138_x_at | io sapiens JTV1 gene (JTV1), mRNA. /FEA=mRNA /GEN=JTV1 /PROD=JTV1 /DB : gb:U24169.1 gb:BC002853.1 |
| | io sapiens hypothetical protein FLJ13287 (FLJ13287), mRNA: /FEA-mRNA /GEN=FLJ13 EF-gi:13375888 /UG-Hs.53263 hypothetical protein FLJ13287 /FL-gb:NM 024647.1 |
| 219714_s_at | gb:NM_018398 1 /DEF=Nomo sapiens calcium channel alpha2-delta3 subunit (HGA272268), mRNA. /FEA=mRNA /GEN=HSA2 /2268 / PROD=calcium channel alpha2-delta3 subunit /DB XREF=g1:8923764 /UG=Hs.22958 calcium channel, voltage-dependent, alpha 2delta 3 |

| | subunit /FL-gb:NM_018398.1 |
|-------------|--|
| 201263_at | gb:NM 003191.1 /DEE=Homo sapiens threonyl-tRNA synthetase (TARS), mRNA. /FEA=mRNA /GEN=TARS /PROD=threonyl-tRNA synthetase / DE XREF=gi:4507366 /UG=HS.84131 threonyl-tRNA synthetase /FL=gb:BC000517.1 qb:M63180.1 qb:NM 003191 1 |
| 201157_s_at | 020500.1 /DEF-Homo sapiens myristoyl CoA:protein N-myristoyltra in N-myristoyltransferase /DB_XREF-g1:2760893 /UG-Hs.111039 N-n 20500.1 gb:AF043324.1 |
| 221593_s_at | gb:BC001663.1 /DEE-Homo sapiens, Similar to ribosomal protein 131, clone MGC:1641, mRNA, complete cds. /FEA-mRNA / PROD-Similar to ribosomal protein 131 /DB XREF-qi:12804504 /UG-Hg 164170 vaccular Rah-Gapmac-cortsining /FIA-mRNA |
| 214202_at | Consensus includes gb:N21364 /FEA=EST /DB_XREF=gi:1126534 /DB_XREF=est:yx50h10.s1 /CLONE=IMAGE:265219 /UG=HS.26968 Homo sapiens mRNA from chromosome 5q21-22, clone:357Ex |
| 203142_s_at | gb:NN 003664.1 /DEF=Homo sapiens adaptor-related protein complex 3, beta 1 subunit (AP3B1), mRNA. /FER=nRNA. /GEN=RP3B1 /PROD=adaptor-related protein complex 3, beta 1subunit /DB XREF=gi:4501974 /UG-Hs.155172 adaptor-related protein complex 3, bera 1 subunit /r=ab.no.031 ich.no.com |
| 222001_x_at | 160126 /FEA=EST /DB_XREF=c |
| 213374_z_at | Consensus includes gb:AW000964 /FEA=EST /DB_XREF=g1:5847880 /DB_XREF=est:wr90h10.x1 /CLONE=IMAGE:2495011 /UG=Hs.236642 3-hydroxyisobutyryl-Coenzyme A hydrolase |
| 201018_at | 1 /DEF-Homo sapiens mRNA full length insert cDNA clone EUROIMAGE 39515. / eukarvotic translation initiation farter la /Fr_dh. DCNATO3 1 ch. 19050 1 |
| 206206_at | IT O |
| 209566_at | Journey 17 June Antigen 64 (mouse) homolog, radioprotective, 105kD /FI-gb:D83597.1 gb:NM 005582.1 Consensus includes gp.H.DEDF=Homo sapiens mRNA; cDNA DKFE64340071 (from clone DKF2p4340071). /FEA=mRNA / DB XREF==1:5267661 (NG=Hs 7089 includes included includes include |
| 212194_s_at | 418892 /FEA=EST / DB XREF=gi:42648 |
| 204526_s_at | |
| 201568_at | gb:NM 014402.1 /DEE=Homo sapiens low molecular mass ubiquinone-binding protein (9.5kD) (QP-C), mRNA. /FEA=mRNA /GEN=CP-C / PRODE_Low molecular mass ubiquinone-binding protein /DB XEEF=gi:7657485 /UG-Hs.3709 low molecular mass ubiquinone-binding protein (9.5kD) /FL=gb:BC001390.1 qb:D50369.1 qb:NM 01402.1 |
| 221829_s_at | Consensus includes gb:AI307759 /FEA=EST /DB_XREF=g1:4002363 /DB_XREF=est:tb24g08.x1 /CLONE=IMAGE:2055326 /UG-Hs.168075 Karyopherin (importin) beta 2 |
| 200613_at | اعاة عا |
| 213160_at | 36964. E cyto |
| 211375_s_at | gb:AF141870.1 /DEF=Homo sapiens translational control protein 80 mRNA, complete cds. /FEA=mRNA /PROD=translational control protein 80 /DB XREF=gi:5006601 /UG-Hs.256583 interleukin enhancer binding factor 3. 90pt /FF=mb.aF141870 1 |
| 202276_at | no sapiens Deleted in split-handsplit-foot 1 region (Iregion /DB_XREF=gi:5453639 /UG=Hs.85215 Deleted in sp |
| 201892_s_at | gb:NN 000884.1 /DEF=Homo sapiens IMP (inosine monophosphate) dehydrogenase 2 (IMPDH2), mRNA. /FEA=mRNA /GEN=IMPDH2 /PROD=IMP (inosine monophosphate) dehydrogenase 2 /DE_XREF=gi:4504688 /UG=Hs.75432 IMP (inosine monophosphate) dehydrogenase 2 / FL=gb:J04208.1 gb:NM 000884.1 |
| 217905_at | gb:NM 024834.1 /DEF=Homo sapiens hypothetical protein FLJ13081 (FLJ13081), mRNA. /FEA-mRNA /GEN-FLJ13081 /PROD-hypothetical protein FLJ13081 /FLJ13081 /PR-PLJ13081 /PR-PLJ130 |
| 209036_s_at | gb:BC001917.1 /DEF=Homo sapiens, malate dehydrogenase 2, NAD (mitochondrial), clone MGC:3559, mRNA, complete cds. / FEA-mRNA /PROD-malate dehydrogenase 2, NAD (mitochondrial) /DB.XREF=gi:12804928 /UG=Hs.111076 malate dehydrogenase 2, NAD (mitochondrial) /FIF=gb:BC001917.1 gb:AF047470.1 gb:NM 005918.1 |
| | |

| 222010_at | Consensus includes gb:BF224073 /FEA-EST /DB XREF-gi:11131299 /DB XREF-est:7q83e05.x1 /CLONE-IMAGE:3704936 /UG-Hs.278544 acetyl-Coenzyme A acetyl-coenzyme A acetylcransferase 2 (acetoacetyl Coenzyme A thiolase) |
|--------------------|---|
| 203825_at | sapiens bromodomain-containing 3 (BRD3), mRNA, /FE. 408642 /UG-Hs.86896 bromodomain-containing 3 /FL-gb |
| | /DEF-Human alpha-N-acetylgalactosaminidase mRNA, complete c 89054 /UG-Hs.75372 N-acetylgalactosaminidase, alpha- /FL-gb |
| | ranscription factor NFATx4 mRNA, complete cds. /FEA-mRNA Hs.172674 nuclear factor of activated T-cells, cytoplasmi |
| | sapiens hypothetical protein FLJ12903 (FLJ12903), mRNA. /FEA-mRNA /GEN-FLJ12903 /PROD-hypothe F-gi:12232418 /UG-HS.14928 hypothetical protein FLJ12903 /FL-gb:NM 022753.1 |
| 208070_s_at | gb:NM 002912.1 /DEE-Homo sapiens REV3 (yeast homolog)-like, catalytic subunit of DNA polymerase zeta (REV3L), mRNA. /FEA-mRNA / GEM-REV3L /PROD=REV3 (yeast homolog)-like, catalytic subunit ofDNA polymerase zeta /DB_XREF=gi:4506482 /UG=Hs.115521 REV3 (yeast homolog)-like, catalytic subunit of DNA polymerase zeta /FL-qb:AF078695.1 gb:MM 002912.1 gb:AF179428.1 |
| 201272_at | sapiens aldo-keto reductase family 1, member B1 (aldose eductase family 1, member B1 (aldosereductase) /DB XREFegose reductase) /FL-gb:BC000260.1 gb:BC005387.1 gb:J04795 |
| 209824_s_at | /DEF-Homo sapiens mRNA for BFALlb, complete cds. /FEA-mRNA /PROD-EWALlb /DB_XREF-gi: ceptor nuclear translocator-like /FL-gb:AB000812.1 gb:AF044288.1 |
| | sapiens mRNA; cDNA DKFZp586C051 (from clone DKFZp586C051); complete cds. /FEA-mRNA /GEN-DKFZp n /DB XREF-gi:12053334 /UG-Hs.109643 polyadenylate binding protein-interacting protein 1 /FL-g |
| at | gb:NN 003204.1 /DEF=Homo sapiens nuclear factor (erythroid-derived 2)-like 1 (NFE2L1), mRNA. /FEA=mRNA /GEN=NFE2L1 / PROD= transcription factor 11 (basic leucine zippertype) /DB_XREF=g1:4505378 /UG=Hs.83469 nuclear factor (erythroid-derived 2)-like 1 / FL=gb:NN 003204.1 gb:U08853.1 |
| 221787_at | 431618 /FEA-EST /DB_XREF-gi:11443732 /DB_XREF-est:7 s clone 24538 mRNA sequence |
| 211581 <u>z</u> at | mRNA, cLST1E splice variant, complete cds. /FEA-mRNA /GEW=LST1 mphocyte antigen 117 /FL=gb:AF000426.1 |
| 37384_at | an mRNA for KIAA0015 gene, com |
| 218423_x_at | |
| 210213_s_at | /FEA=mRNA / ng protein / |
| 201598_s_at | 40.00 |
| 204301_at | |
| 202877_s_at | Consensus includes gb:W72082 /FEA=EST /DB_XREF=gi:1382588 /DB_XREF=est:zd70c06.sl /CLONE=IWAGE:345994 /UG=Hs.97199 complement component Clg receptor /FL=gb:NM 012072.2 gb:U94333.1 |
| 208772_at | /DB_XREF=gi:11022197 /DB_XREF=est:AU160676 / |
| 201998_at | Consensus includes gb.AI743792 /FEA-EST /DB XREF-gi:5112080 /DB XREF-est:wg53h11.x1 /CLONE-IMAGE:2368869 /UG-Hs.2554 sialyltransferase 1 (beta-galactoside alpha-2,6-sialytransferase) /FL-gb:NM 003032.1 |
| 208722_s_at | sapiens complex 1950.1 |
| 55692_at | |
| 217954_s_at | gb:NM 015153.1 /DEF-Homo sapiens KIAA0244 protein (KIAA0244), mRNA. /FEA-mRNA /GEN-KIAA0244 /PROD-KIAA0244 protein / DB XREF-gi:7662017 /UG-Hs.78893 KIAA0244 protein /FL-gb:AF091622.1 gb:NN 015153.1 |

Table 2: Genes from Cluster Analysis 2

| Affymetrix-internal designation | Description of the sequence in the GeneBank data base |
|---------------------------------|--|
| 200654_at | gb:302783.1 /DEF-Human thyroid hormone binding protein (p55) mRNA, complete cds. /FEA-mRNA /GEN-P44B /DB_XREF-gi:339646 /UG-Hs.75655procollagen-proline, 2-oxoglutarate 4-dioxygenase (proline 4-hydrozylase), hera nolyoeptide (protein disulfide isomerase; thyroid hormone bindingprotein p55) /FL-9b:J02783.1 gb:NM 900918.1 |
| 203126_at | gb:NM 014214.1 /DEF-Homo saplens inositol(myo)-1(or 4)-monophosphatase 2 (IMPA2), mRNA. /FEA-mRNA /GEN-IMPA2 / ppo-inositol(myo)-1(or 4)-monophosphatase 2 /DB_XREF-gi:7657235 /UG-Hs.5753 inositol(myo)-1(or 4)-monophosphatase 2 //FL-ob:AF200432.1qb:NN 014214.1 gb:AF014398.2 |
| 203585_at | SD:NM 007150.1 /DEF-Homo sapiens zinc finger protein 185 (LIM domain) (ZNF185), mRNA. /FEA-mRNA /GEN-ZNF185 - (DIM 007150.1 / PEA-MRNA / PEA-MR |
| 205220_at | gb:NM 006018.1 /DEF=Homo sapiens putative chemokine receptor; GTP-binding protein (HM4), mkNA. /кем-mким /GEN-HN74 /PROD-putative chemokine receptor; GTP-bindingprotein /DB_XREF=gi:5174460 /UG-Hs.137555 putative chemokine receptor; GTP-binding protein /FL-gb:NM 006018.1 |
| | db:NN 001557.1 /DEF-Homo sapiens interleukin 8 receptor, beta (ILBRB), mRNA. /FEA-mRNA /GEN-ILBRB /FROD-interleukin 8 receptor, beta /FL-9b:N94582.1 gb:NJ3469.1 gb:NJ 001557.1 db:L19593.1 beta /DB XREF-91:4504682 /UG-HS.846 interleukin 8 receptor, beta /TRA /FL-9b:N94582.1 gb:NJ 001557.1 db:L19593.1 |
| at | gb:NR 016081.1 /DEF-Homo sapiens palledin (KIAMU994), MRNA. /ELA-MRNA /GEN-ALANASZZ / KNOC-Palledin / EL-MINISTER / DEF-HOMO Sapiens / DEF-HOMO SAPER / DEF-HOMO SAPER / DEF-HOMO / DEF-HOME / DEF-HOM |
| 205568_at | qb:lNM 020980.2 /DEF-Homo sapiens aquaporin 9 (AQP9), mKNN. /LEA-mKNN /SEN-AQE9 /FROD-aquaporin 9 /FI-gb:NN 020980.2 gb:AB008775.1 gb:AF016495.1 UG-Hs.104624 aquaporin 9 /FI-gb:NN 020980.2 gb:AB008775.1 gb:AF016495.1 |
| 214022_s_at | Consensus includes gb:AA749101 /FEA-EST /DB_XREF-gi:2/89059 /Db_XREF-est:nyilouz.si /LLONE-inaus.iz.iz./ / UG-Hs.146360 interferon induced transmembrane protein 1 (9-27) |
| 209762_x_at | gb:AF280094.1 /DEE=Homo sapiens transcriptional coactivator SpilDb mRNA, complete cds. / FLA=mRNA. / 52kD /FL=gb:AF280094.1 /PROD=transcriptional coactivator SpilOb /DB XREF=g1:9800493 /UG-Hs.38125 interferon-induced in 75, 52kD /FL=gb:AF280094.1 / PROD=transcriptional coactivator SpilOb /DB XREF=g1:9800493 /UG-Hs.38125 interferon-induced in 75, 52kD /FL=gb:AF280094.1 |
| 203234_at | gb:NM 003364.1 /DEF-Homo sapiens uridine phosphorylase (UP), MKNA. /EEN-MONA /GEN-OF /FNOL-ULIQUINE phosphorylase /EL-gb:EC001405.1 gb:NM 003364.1 /OB XREF-gi:4507838 /UG-Hs.77573 uridine phosphorylase /FL-gb:EC001405.1 gb:NM 003364.1 /OF AREF-gi:4507838 /UG-Hs.77573 uridine phosphorylase /FL-gb:EC01405.1 gb:NM 003364.1 /UG-Hs.77573 uridine / |
| 201940_at | Consensus includes gb:AA897514 /FEA=EST /DB KKEF=g1:U34134 /DB AKEF=EST:3J02D1Z:31 /LDDRE_IENGE:1539015 / CONSENSUS 1559015 / |
| 209310_s_at | academ zere |
| <u>202128_at</u> | gb:NM 014821.1 /DEE-Homo sapiens KLAAU11/ gene product (hizaU211/) mkWh. /rrAmwa /srkminan/1 db:NM 014821.1 //PROD-KIAA0317 gene product /DB XREF-g1:7662051 /UG-Hs.20126 KIAA0317 gene product /FL-gb.201215.1 gb:NM 014821.1 |
| 201192_s_at | gb:NM 006224.1 /DEE=Homo sapiens phosphotioyiinositoi transier piotein (Filen), many. /ren-many /och.iii., / PROD-phosphotidylinositol transfer protein /DB_XREF-gi:5453907 /UG-Hs.79709 phosphotidylinositol transfer protein / FL-gb:D30036.1 gb:NM3704.1 gb:NM 006224.1 |
| 208012_x_at | gb:NM 004509.1 /OEF=Homo sapiens interferon-induced protein 41, 30kU (11141), mkNA: /FELT-mANA /VENT-1111 / PROD-Interferon-induced protein 41, 30kD /DB_XREF=g1:4758585 /UG-Hs.241510 interferon-induced protein 41, 30kD / FL-gb:122342.1 gb:NM 004509.1 |
| 209137_s_at | gb:BC000263.1 /DEF-Homo sapiens, Similar to ubiquintin c-terminal hydrolase related polypeptide, cloud not.zozi, mnnn, complete cds. /FEA-mRNA /PROD-Similar to ubiquintin c-terminal hydrolaserelated polypeptide /DB_XREF-g1:12653004 / UG-Hs.78829 ubiquitin specific protease 10 /FL-gb:BC000263.1 |
| 218023_s_at | gb:RM 016605.1 /DEF-Homo sapiens putative nuclear protein (LUC130.1), mRNA: /FEA-MRNA /UEN-LUC1315.7 / PROD-putative nuclear protein /DB XREF-gi-1706138 /UG-H2 Putative nuclear protein /FE-gb:RF251040.1 gb:NM 016605.1 |
| 201190_s_at | Consensus includes gb:Hlb64/ /EEA=E31 /UB AKER=G3:80040/ /UB AKER=G3L.7M2/003121 / CLONE 11005221.1 UG-H3.79709 phosphotidylinositol transfer protein /Firefsiphone for minimal complete / FFB=mpNB / |
| 210563_x_at | db:U97075.1 /DEF*Homo sapiens FLICE-11Ke innibitory protein short form March, Compiere Cos. / Lon March Febb- PROD-FLICE-11Ke inhibitory protein short form /DB XREF=gi:2253680 /UG-Hs:195175 CASP8 and FADD-11ke apoptosis regulator / |
| | |

| | EL=gb:U97075.1 |
|-------------|---|
| 221827_at | XREF-gi:10209637 /DB_XR |
| 202041_s_at | gb:NM 004214.3 /DEF-Homo sapiens fibroblast growth factor (acidic) intracellular binding protein (FIBP), mRNA. / FER-mRNA /GENFIRD /PROD-fibroblast growth factor (acidic) intracellularbinding protein /DB XREF-gi:7262377 / 10-HB.7768 fibroblast growth factor (acidic) intracellular binding protein /FL-gb:AF171944.I gb:AF171945.1 gb:AF171946.1 9b:AF1010187.2 gb:NN 004214.3 |
| 201001_s_at | Consansus includes gb:BG164064 /FEA=EST /DE XREF=gi:12670767 /DB XREF=est:602341091F1 /CLONE=INAGE:4449022 / UG-Hs.75875 ubiquitin-conjugating enzyme E2 variant 1 /FL-gb:U39361.1 qb:NM 003349.2 qb:EC000468.1 |
| 212975_at | |
| 201412_at | 12 7 |
| 203708_at | 9b:NM 002600.1 /DEF=Homo sapiens phosphodiesterase 4B, CAMP-specific (dunce (Drosophila)-homolog phosphodiesterase E4 (PDE4B) mRNA. /FEA=mRNA /GEN=PDE4B /PROD=phosphodiesterase 4B, CAMP-specific (dunce Drosophila)-homolog phosphodiesterase E4 (PDE4B) |
| | E-g1:4505662 /UG-H3:188 phosphodiesterase 4B, cAMP-specific (dunce (brosophila)-homolog phosphodiesterase E4) |
| 203907_s_at | gb:NM 014869.1 /DEF-Homo sapiens KIAA0763 gene product (KIAA0763), mRNA. /FEA-mRNA /GEN-KIAA0763 /PROD-KIAA0763 gene product / DB XREF-gi:1662289 /UG-Ns.4764 KIAA0763 gene product /FL-gb:AB018306.1 gb:NM 014869.1 |
| 209882_at | gb:AF084462.1 /DEF-Homo sapiens GTP-binding protein ROC1 (ROC1) mRNA, complete cds. /FEA-mRNA /GEN-FOC1 / PROD-GTP-binding protein ROC1 /DB XREF-gi:4234917 /UG-Hs.96038 Ric (Drosophila)-like, expressed in many tissues / FL-gb:U71203.1 qb:U78165.1 qb:AF084462.1 qb:NM 006912.1 |
| 204276_at | Consensus includes gb:BE895437 /FEA-EST /DB XREF-gi:10358829 /DB XREF-est:601437912F1 /CLONE-INAGE:3922971 / UG-Hs.274701 thymidine kinase 2; mitochondrial /FL-gb:NM 004614.I gb:U77088.1 |
| 221984_s_at | des gb:AL040896 / othetical protein |
| 218517_at | gb:NM 024900.1 /DEF=Homo sapiens hypothetical protein FLJ22419 (FLJ22419), mRNA. /FEA=mRNA /GEN=FLJ22419 / PROD-hypothetical protein FLJ22419 /DB XREF-gf:13376356 /UG-Hs.238246 hypothetical protein FLJ22419 /FL-qb:NN 024900.1 |
| 208740_at | Consensus includes gb:BF593650 /FEA=EST /DB XREF=gi:11685974 /DB XREF=est:nac02d03.x1 /CLONE=IMAGE:3275957 / UG=Hs.23964 sin3-associated polypeptide, 18kD /FI=gb:NM 005870.2 gb:U96915.1 gb:AF153608.1 gb:U78303.1 |
| 221484_at | Consensus includes gb:BF691447 /FEA-EST /DB XREF=gi:11976855 /DB XREF=est:602247615F1 /CLONG-INAGE:4332866 / UG-Hs:107526 UDP-Gal:betaGlcNAc beta 1,4- galactosyltransferase, polypeptide 5 /FL-db:AB004550.1 db:AF038663.1 db:NN 004776.1 |
| 209575_at | no sapiens, Similar to interleukin 10 receptor, beta, clone MGC:2210, mRNA, complete cds. r to interleukin 10 receptor, beta /DB_XREF=gi:12804902 /UG-Hs.173936 interleukin 10 recept gb:NM 000628.1 |
| 200619_at | |
| 217286_s_at | Consensus includes gb:BC001805.1 /DEF-Homo sapiens, clone IMAGE:3543670, mRNA, partial cds. /FEA-mRNA / PROD-Mixnown (protein for IMAGE:3543670) /DB_XREF-gi:12804742 /UG-Hs.240615 hypothetical protein FLJ13556 similar to N-myc |
| 207842_s_at | gb:NM 007359.1 /DEF=Homo sapiens MLN51 protein (MLN51), mRNA. /EEA-mRNA /GEN-NLN51 /PROD-MLN51 protein / DB XREF-gi:6678887 /UG-Hs.83422 MLN51 protein /FL-qb:NM 007359.1 |
| 217788_s_at | <pre>dpiNM 004481.2 /DEF=Homo sapiens UDP-N-acetyl-alpha-D-galactosamine:polypeptide N-acetylgalactosaminyltransferase 2 (GalNAc-T2) (GALNT2), mRNA. /FEA-mRNA /GEN-GALNT2 /PROD-polypeptide N-acetylgalactosaminyltransferase 2 / DB XREF=qi:9945385 /UG=Hs.130181 UDP-N-acetyl-alpha-D-galactosamine:polypeptide N-acetylgalactosaminyltransferase 2</pre> |
| 48612_at | 1. AA225490 |
| 212689_s_at | Consensus includes gb:AA524505 /FEA-EST /DB_XREF-gi:2265433 /DB_XREF-est:ng43gl2.sl /CLONE-IMAGE:937606 / UG-Hs.321707 KIAA0742 protein |

| 222047_s_at | Consensus includes gb:AI523895 /FEA=EST /DB XREF=g1:4438030 /DB_XREF=est:tg97g03.x1 /CLONE=IMAGE:2116756 / UG-Hs.111801 arsenate resistance protein ARS2 |
|----------------------|---|
| 201666_at | omo sapiens tissue inhibitor of metalloproteinase 1 (erythroi RNA. /FEA-mRNA /GEN-TIMP1 /PROD-tissue inhibitor of metallop G-Hs.5831 tissue inhibitor of metalloproteinase 1 (erythroid 00866.1 gb:M12670.1 gb:M59906.1 gb:NM 003254.1 |
| 203218_at | . 037431 /FEA-EST /DB_XREF-gi:1319025 /DB_XREF-est:zcllf11.s1 /CLONE-INAGE:322029 / activated protein kinase 9 /FL-gb:U34821.1 gb:MM 002752.1 gb:L31951.1 gb:U09759.1 |
| 213118_at | Consensus includes gb:AL136821.1 /DEF=Homo sapiens mRNA; cDNA DKF2p434M1526 (from clone DKF2p434M1526). /FEA=mRNA / GEN-DKF2p434M1526 /PROD-hypothetical protein /DB XREF=gi:12053152 /UG-Hs.153293 KIAA0701 protein |
| 201057_s_at | <pre>db:NM 004487.1 /DEE=Homo sapiens golgi autoantigen, golgin subfamily b, macrogolgin (with transmembrane signal), 1 (GOLGB1), mRNN. /FEA=MRNA /GEN=GOLGB1 /PROD=golgi autoantigen, golgin subfamily b, macrogolgin (with transmembrane signal), 1 / DB_XREF=gi:4758653 /UG=Hs.7844 golgi autoantigen, golgin subfamily b, macrogolgin (with transmembrane signal), 1 / EL=gp:NN 004487.1.</pre> |
| 217738_at | :BF575514 /FEA=E9 |
| 208648_at | :W60953 /FEA=EST /DB XREF=gi:1367731 /DB XREF=est:zc98bl2.s1 /CLONE-IMAGE:339167 / containing protein /FL=gb:AF100752:1:qb:NM 007126:2 |
| 212242_at | /FEA-EST /DB_XREF-gi:12916087 /DB_XREF-est:AL565074 /CI estis specific) |
| 203897_at | FEA-EST /DB_XREF-g1:11766863 /DB_X A-211C6.1 /FL-gb:NM 020424.1 |
| 209513_s_at | Similar to RIKEN cDNA 2610207116 gene, clone MGC:10940, mRNA, complete cds. / cDNA 2610207116 gene 70B XREF=g1:13279253 /UG=Hs:47986 Homo sapiens, gene, clone MGC:10940, mRNA, complete cds /FL=gb:BC004331.1 |
| 211762_s_at | nmo sapiens, karyopherin alpha 2 (RAG cohort 1, importin alpha 1), clone MGC: herin alpha 2 (RAG cohort 1, importinalpha 1) /DB XREF-gi:13543656 /FL-gb:BC |
| 219639 <u>-</u> x_at | lomo sapiens pothetical p 16 /FL=gb:NM |
| 217882_at | omo sapiens 30 kDa protein (LOC55831), mRNA. /FEA-mRNA /GEN-LOC55831 /FROD-30 kDa prot G-Hs.283714 30 kDa protein /FL-gb:AF157321:1 gb:NM 018447.1 |
| 204714_s_at | o sapiens coagulation factor V (proaccelerin, labile factor) (F5), mRNA: /FEA=mRNY V precursor /DB_XREE=gi:10518500 /UG=Hs.30054 coagulation factor V (proaccelerin, 5967.1 gb:M14335.1 |
| 221547_at | mo sapiens, pre-mRNA splicing factor similar to S. cerevisiae Prp18, clone MGC:5075, NA /PROD-pre-mRNA splicing factor similar to S.cerevisiae Prp18 /DB_XREF-gi:12653992 , processing factor 18 /FL-gb:BC000794.1 gb:U51990.1 gb:NM.003675.1 |
| 210793_s_at | an nucleoporin 98 (NUP98) mRNA, complete cds. /FEA-mRNA /GEN-NUP98 NG-Hs.112255 nucleoporin 98kD//EL-gb:U41815.1 |
| 206207_at | lomo sapiens Charot-Leyden crystal protein (CLC), mRNA. /FEA-mRN. ystal protein /DB_XREF-qi:6325464 /UG-Hs.889 Charot-Leyden crys 001828:3 |
| 202595_s_at | mo sapiens HSPC112 mRNA, complète cds. /FEA-mRNA /PROD-HSPC112 /DB_XREF=g1:6841445 / ceptor overlapping transcript-like 1 /FL-gb:BC000642.1 gb:AF063605.1 gb:AF161461.1 gb:A |
| 219434_at | lomo sapiens triggering receptor expressed on myeloid cells 1 (TREM) vtor expressed on myeloid cells 1 /DB XREF=g1:8924261 /UG=Hs.283022 th:AF196329.1 gb:NM 018643.1 gb:AF287008.1 |
| 205020_s_at | <pre>lomo sapiens ADP-ribosylation factor-like 4 (ARL4), mRNA, /FEA=mRNA /GE i factor-like 4 /DB_XREF-g1:5031602 /UG=Hs.201672 ADP-ribosylation fact 005738.1</pre> |
| 212052_s_at | Consensus includes gb:AB014576.1 /DEF-Homo sepiens mRNA for KIAA0676 protein, partial cds. /FEA-mRNA / GEN-KIAA0676 /PROD-KIAA0676 protein /DB XREF-gl:3327165 /UG-Hs.155829 KIAA0676 protein |

| 202565_s_at | gb:NM_003174.2 /DEF-Homo sapiens supervillin (SVIL), transcript variant 1, mRNA. /FEA-mRNA /GEN-SVIL / PROD-supervillin, isoform 1 /DB XREF-gi:11496980 /UG-Hs.154567 supervillin /FL-qb:NM 003174.2 db:AF051850.1 db:AF051851.1 | _ |
|---------------------|--|------------|
| 221524_s_at | omplete cds. /FEA=mRNA /PROD=R.gb:AF272036.1 gb:BC003088.1 | |
| 218037_at | gb:NM 024293.1 /DEF-Homo sapiens hypothetical protein MGC3035 (MGC3035), mRNA. /FER-mRNA /GEN-MGC3035 / PROD-hypothetical protein MGC3035 / FROD-hypothetical protein MGC3035 / FROD FROD FROD FROD FROD FROD FROD FROD | · . |
| 214107_x_at | Consensus includes gb:AW340850 /FEA=EST /DB XREF=g1:6837476 /DB XREF=est:xx31f01.xl /CLONE=TNAGE:2761753 /UG-Hs.326350 Homo sapiens cDNA FLJ11822 fis, clone HEMBA1006485, highly similar to PUROMYCIN-SENSITIVE ANINOPEPTIDASE (EC 3.4.11) | ; ; |
| 210184_at | adhesion glycor in, alpha X (an | |
| 209107_x_at | . /FEA-mRNA /GEN-Hin-2 :U19179.1 | |
| 208749_x_at | <pre>gb:AF085357.1 /DEF=Homo sapiens flotillin mRNA, complete cds. /FEA=mRNA /PROD=flotillin /DB_XREF=g1:5114048 / UG=Hs.179986 flotillin 1 /FL=gb:BC001146.1 gb:AF085357.1 gb:AF089750.2 qb:NN 005803.2</pre> | |
| 212561_at | | |
| 212470_at | Consensus includes gb:AB011088.1 /DEF=Homo sapiens mRNA for KIAA0516 protein, partial cds. /FEA=mRNA /GEN=KIAA0516 / PROD=KIAA0516 protein /DB XREF=gi:3043555 /UG-Hs.129872 sperm associated antigen 9 | |
| 214937_x_at | Consensus includes gb:AI924817 /FEA=EST /DB_XREF=gi:5660781 /DB_XREF=est:wn23c11.xl /CLONE=IMAGE:2446292 / UG=Hs.75737 pericentriolar material 1 | <u>.</u> . |
| 201783 <u>_s</u> at | otheliosis viral onco (RELA), mRNA. /FEA olog A (nuclear facto avian reticuloendoth | |
| 202010_s_at | INUCLEAR LACTOL OF KAPPA .1.90T POLYPEPLINE Gene enhancer in B-Cells 3 (PbS)) /FL=4p:NM 021975.1 qb:L19067.1 qb:LVM 021881.1 /DEF=HOMO sapiens clones 23667 and 23775 zinc finger protein (LOCS7062), mRNN. /FER=mRNN /GEN=LOCS7062 / FROM=LOCS7062 / FROM=LOC | |
| . at | gb:NM 003641.1 /DEF=Homo sapiens interferon induced transmembrane protein 1 (9-27) (IFITM1), mRNA. /FEA=mRNA /GEN=IFITM1 / PROD=Interferon induced transmembrane protein 1(9-27) /DB XREF=gi:4504580 /UG=Hs.146360 interferon induced transmembrane protein 1 (9-27) /FI=qb:BC000897,1 qb:J04164.1 qb:NM 003641.1 | |
| 203266_s_at | gb:NM 003010.1 /DEF-Homo sapiens mitogen-activated protein kinase kinase 4 (NMP2K4), mRNA. /EEA-mRNA /GEN-MAP2K4 / PROD-mitogen-activated protein kinase kinase 4 /DB_XREF-gi:4506888 /UG-Hs.75217 mitogen-activated protein kinase kinase 4 / FEL-gb:RN 003010.1 gb:136870.1 gb:U17743.1 | : ' ' |
| 219863 <u>a</u> t | gb:NM 016323.1 /DEE-Homo sapiens cyclin-E binding protein 1 (LOC51191), mRNA. /FEA-mRNA /GEN-LOC51191 / PROD-cyclin-E binding protein 1 /DB XREF-qi:7705930 /UG-Hs.26663 cyclin-E binding protein 1 /FL-qb.AB027289.1 db:NM 016323.1 | • |
| 212322_at | REF-est:7h15b02.x1 /CLONB-IMAGE:3316011./ | |
| 204206_at | gb:NM 020310.1 /DEE-Homo sapiens MAX binding protein (MNT), mRNA. /FEA-mRNA /GEN-MNT /PROD-MAX binding protein / DB XREF-qi:9945317 /UG-Hs.25497 MAX binding protein /FI-qb:NM 020310.1 | |
| 203278_s_at | gb:NM 016621.1 /DEF-Homo sapiens hypothetical protein (LOC51317), mRNA. /FERA-mRNA /GEN-LOC51317 / PROD-hypothetical protein /DB XREF-gi:7706159 /UG-Ms.106826 KIAA1696 protein /FL-gb:AF208848.1 gb:NM 016621.1 | . • • |
| 206637_at | ne product; putative G-protein-coupled receptor; G /GEN=KIAA0001 /PROD=KIAA0001 gene product; putativ glane1:17661847 /UG-Hs.2465 KIAA0001 gene product glucose /EL-gb:D13626.1 gb:NN 014879.1 | * : |
| 18924_s_at | gb:NM_004388.1 /DEF-Homo sapiens chitcbiase, di-N-acetyl- (CTBS), mRNA. /FEA-mRNA /GEN-CTBS /PROD-chitcbiase, di-N-acetyl- /DB XREF-gi:4758091 /UG-Ms.135578 chitcbiase, di-N-acetyl- /FL-gb:M95767.1 gb:NM 004388.1 | |

Table 3: Genes from Cluster Analysis 3

| Affvmetrix | Description of the sequence in the GeneBank data base |
|-------------|--|
| internal | |
| designation | Charer Incl. A1021977; bK441C4.1 (novel NAFF (v-maf musculoaponeurotic fibrosarcoma (avian) oncogene family, protein F) |
| 78 77,00 | protein) /cds=(0,494) /gb=AL021977 /gi=4914526 /ug=Hs.51305 /len=2128 protein) /cds=(0,494) /gb=AL021977 /gi=4914526 /ug=Hs.51305 /len=2128 |
| 210845_s_at | gb:U08839.1 /Dbt=Human Urokinase-Lype Fical 1957 UG-Hs.179657 plasminogen activator, urokinase receptor /FL-gb:U08839.1 plasminogen activator receptor /DB XREF-gi:517197 /UG-Hs.179657 plasminogen activator, urokinase receptor /DB XREF-gi:517197 /UG-Hs.17960 tumor |
| 202643_s_at | Consensus includes gb:Al738896 /FEA-EST /DB KKEr91:31008/ /DB KKEr91:32008/ /DB KKER |
| 205476_at | gb:NM 004591.1 /DEF-Homo sapiens small inductble cytokine subtainty A (cys-Cys). Minimization 1759015 /UG-Hs.75498 small inductble CRP-SCAPI inductble cytokine subfamily A (cys-Cys), member 20 /DB XREF-1590015 /UG-Hs.75498 small inductble cytokine subfamily A (cys-Cys), member 20 /EL-ab:UG-Hs.75498 small inductble cytokine 20 /EL-ab:UG-Hs.77035.1 gb:DB6955.1 gb:NM 004591.1 |
| 205419_at | Cytokine Supremary. Option Captern Special Epstein-Barr virus induced gene 2 (lymphocyte-specific G protein-coupled receptor) (Ebil), app. M. Option Coupled receptor) / Epstein-Barr virus induced gene 2 (lymphocyte-specific G protein-coupled receptor) / MRNA (GRI=EB12 /PROD-Epstein-Barr virus induced gene 2 (lymphocyte-specific G protein-coupled receptor) / |
| | DB XRE-gi:4826705 /UG-Ns.784 Epstein-Barr virus induced gene 2 (Lymphocyte-specific & protesti-Coupied receptor) / Epstein-Barr virus induced gene 2 (Lymphocyte-specific & protestiv acrivation antigen CD69 / |
| 209195_at | gb:107555.1 /DEE-Homo sapiens early activation antigen CD69 mRNA, complete cus. /FEA-WRNA / 10555.1 db:NN 001781.1 DB XREF-q1:291897 /UG-Hs.82401 CD69 antigen (p60, early T-cell activation /FE-dp:107555.1 db:NN 001781.1 |
| 205767_at | gb:NN 001432.1 /DEF=Homo sapiens epiregulin (EREG), mRNA. /FEA=mRNA /GEN=EREG /FNOU=epiregulin processor, / |
| 203821_at | OGENS. 113733 EPITEGRAFIA STORM AND TRANSPORT (NEDATIN-binding epidermal growth factor-like growth factor) (DIK), gb:NN 001945.1, DEF-Home Appendent factor) / |
| 1 | mRNA. (FEA-mRNA /GEN=Us /FROD-Olyminatia toxin receptor (heparin-binding epidermal growth factor-like growth factor) / DB XRER-gi:4503412 /UG=Hs.799 diphtheria toxin receptor (heparin-binding epidermal growth factor) |
| | EL-gb: M60278.1 gb: NM 001945.1 |
| 211924_s_at | gb:AY029180.1 / DEF=Homo sapiens soluble utokinase plasminogen activator receptorprecursor / DB XREF=gi:13641308 /FL-gb:AY029180.1 FRA=CDS /GEN=SUPAR / PROD=soluble urokinase plasminogen activator receptorprecursor / DB XREF=gi:13641308 /FL-gb:AY029180.1 |
| 205403_at | gb:NN 004633.1 /DEE-Homo sapiens interleukin 1 receptor, type II (LIRZ), mRNA. /rehmmun /cemminnz /PROD-interleukin 1 receptor, type II /DB_XREE-g1:4758597 /UG-Hs.25333 interleukin 1 receptor, type II |
| | /FL-qb:U74649.1 gb:NM 004633.1 |
| 204351_at | gb:NN U05980.1 /DEr=nomO Septens 3100 Calcium-binding protein P /FL=gb:NN 005980.1 protein P /DB XREF=gi:5174662 /UG-Hs.2962 S100 Calcium-binding protein P /Frammons /Grn=FGR3 /PRODe-early growth response 3 / |
| 206115_at | / grammad/ gram tracas |
| 204103_at | gb:NM 002984.1 /DEF-Homo sapiens small inducible cyfokine A4 (homologous co muuse mip-iul (st.n.), mkkn. |
| 206522_at | -y-cort. /FEA-mRNA /GEN-MGAM / gb:NN 004661 //DEF-Homo sapiens maitase-glucoamylase (alpha-glucoamylase (alpha-glucosidase) /DB_XREF-gl:4158711 /UG-Hs.122785 maitase-glucoamylase (alpha-glucosidase) /DB_XREF-gl:4158711 /UG-Hs.122785 maitase-glucoamylase |
| | /FL-gb:AF016833.1 gb:NM 004058.1 |
| 202147_s_at | gb:kN U01330.1 /DEF-now September 1 /DB XREF-gi:4504606 /UG-Hs.1879 interferon-related developmental PROD-interferon related developmental regulator 1 /DB XREF-gi:4504606 /UG-Hs.1879 interferon-related developmental |
| 206515_at | regulator 1 / L. go nega processor of the part of the polypeptide 3 (leukotriene B4 omega hydroxylase) (CYP4E3), gb:NM 000896.1 / DEF-Home part of the part of the polypeptide 3 / DB XREF-g1:4503240 / UG-Hs.106242 |
| | mkNA. /redemnna /ush-ciris / wolypeptide 3 (leukotriene B4 omega hydroxylase) /FL-gb:AB002454.1 gb:D12620.1 gb:NH 000896.1 cytochrome P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / redemne P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / Redemne P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / Redemne P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / Redemne P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / Redemne P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / Redemne P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / Redemne P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / Redemne P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / Redemne P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / Redemne P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / Redemne P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / Redemne P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / Redemne P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / Redemne P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / Redemne P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / Redemne P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / Redemne P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / Redemne P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / Redemne P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / Redemne P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / Redemne P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / Redemne P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / Redemne P450, subfamily IVF, polypeptide 3 (leukotriene B4 omega hydroxylase) / |
| 204614_at | 9b:NM 002575.1 /DEF=Homo sapiens serine (or cysteine) proteinase inhibitor, cladeB (ovalbumin), member 2 /DB XREF=gi:4505594 / |
| | The state of the s |

| | UG-Hs.75716 serine (or cysteine) proteinase inhibitor, clade B (ovalbumin), member 2 /FL-gb:J02685.1 gb:J03603.1 qb:M18082.1 gb:NM 002575.1 |
|-------------|--|
| 209959_at | |
| 215078_at | Consensus includes 9b:AL050388.1 /DEF-Homo sapiens mRNA; cDNA DKF2p564M2422 (from clone DKF2p564M2422); partial cds. / FEA-mRNA /GEN-DKF2p564M2422 /FROD-hypothetical protein /DB_XREF-g1:4914612 /UG-Hs.306320 Homo sapiens mRNA; cDNA DKF2p564M2422 (from clone DKF2p564M2422); partial cds |
| 201890_at | Consensus includes gb: BE966236 /FEA-EST /DB XREF=gi:11771437 /DB XREF=est:601660172R1 /CLONE=IMACE:3905920 /UG-Hs.75319 ribonucleotide reductase M2 polypeptide /FLT=qb:NN 001034.1 |
| å | E 60 |
| 203888_at | gb:NM 000361.1 /DEF=Homo sapiens thrombomodulin (THBD), mRNA. /FEA=mRNA /GEN=THBD /PROD=thrombomodulin /DB_XREF=g1:4507482 / UG=HS.2030 thrombomodulin /FL=gb:M16552.1 gb:NM 000361.1 |
| 218723_s_at | sapiens RGC32 protein (RGC32), mRNA. /F 1 /FL-qb:AF036549.1 gb:NM 014059.1 |
| 206834_at | gb:NM 000519.2 /DEF=Homo sapiens hemoglobin, delta (HBD), mRNA. /FEA-mRNA /GEN=HBD /PROD-hemoglobin, delta /DB_XREF=g1:6633803 /UG=H5.36977 hemoglobin, delta /FL=gb:NM 000519.2 |
| 205479_s_at | jb:NM 002658.1 /DEF=Homo sapiens plasminogen activator, urokinase (PLAU), mRNA. /FEA-mRNA /GEN=PLAU /PROD-plasminogen activator, urokinase /FL=qb:N15476.1 qb:NN 002658.1 |
| 202912_at | o sapiens adrenomedullin (ADM), mRNA. /FEA-mRNA /GEN-ADM /PROD-adrenc n /FL-gb:NM 001124.1 gb:D14874.1 |
| 208869_s_at | pb:AF087847.1 /DEF-Homo sapiens GABA-A receptor-associated protein like 1 (GABARAPLI) mRNA, complete cds. /FEA-mRNA / GEN-GABARAPLI /PROD-GABA-A receptor-associated protein like 1 /DB XREF-gi:13375570 /UG-HS.282654 Homo sapiens mRNA; cDNA DKF20564N1272 (from clone DKF20564N1272); complete cds /FL-qD:AL136676.1 qb:AF087847.1 |
| 202193_at | sapiens LIM domain kinase 2 (LIMK2) isoform 2a /DB XREF=qi:8051619 /UG= |
| 214696_at | AF070569.1 /DEF-Homo sapiens clone 24659 mRNA sequence. /FEA-mRNA /DE 59 mRNA sequence |
| 38037_at | Cluster Incl. M60278:Human heparin-binding EGF-like growth factor mRNA, complete cds /cds=(261,887) /gb=M60278 /gi=183866 / ug=Hs.799 /len=2342 |
| 212723_at | K021780.1 /FEA=mRNA |
| 207802_at | no sapiens ic granule pas ; cysteine |
| 209967_s_at | mRNA for hCREM (cyclic AMP- CREM-2; hCREM-2 /PROD=hCREM /DB XREP=q:532036 /HG=Hs 1 |
| 210119_at | Inward rectifier potassium channel (Kirl.3), complete cds. /FEA=mRNA /GEN-Kirl.3 /PRN REF=gi:1765984 /UG=Hs.17287 potassium inwardly-rectifying channel, subfamily J, memb?243 1 |
| 203435_s_at | gb:NM 007287.1 / DEE=Homo sapiens membrane metallo-endopeptidase (neutral endopeptidase, enkephalinase, CALLA, CD10) (NME), transcript variant Dis, mRNA, /FEA=mRNA /GEN=MAE /PROD=membrane metallo-endopeptidase, /DB_XREF=gi:6042199 /UG=Hs.1298 membrane metallo-endopeptidase, rangonal and contral endopertidase, enkephalinase, CALLA, CD10) /Fi=mh:103779 i ch:NN 007287 i ch:NN 007287 i |
| 213515_x_at | 1133353 /FEA=EST /DB_XREF=g1:6360669 /DB_XREF=est:HA1957 /VG=Hs.298161 myosin, light polype ic |
| 209545_s_at | gb:AF064824.1 /DEF-Homo sapiens CARD-containing ICE associated kinase mRNA, complete cds. /FEA-mRNA /PROD-CARD-containing ICE associated kinase /DB XREF-gi:3290111 /UG-Hs.103755 receptor-interacting serine-threonine kinase 2 /FL-gb:BC004553.1 gb:AF07406.1 gb:AF064824.1 gb:AF078530.1 gb:AF078530. |
| 212531_at | Consensus includes gb:NM 005564.1 /DEF-Homo sapiens lipocalin 2 (oncogene 24p3) (LCN2), mRNA. /FEA-CDS /GEN-LCN2 / PROD-lipocalin 2 (oncogene 24p3) /DB XREF-gi:5031852 /UG-Hs.204238 lipocalin 2 (oncogene 24p3) /FI-gb:NM 005564.1 |

| 208470_s_at | cDS /GEN=HPR /PROD=haptoglobin-r 1 |
|-------------|---|
| 200733_s_at | gb:U48296.1 /DEF=Homo sapiens protein tyrosine phosphatase FTPCAAX1 (hPTPCAAX1) mRNA, complete cds. /FEA=mRNA /GEN-hPTPCAAX1 / PROD-protein tyrosine phosphatase PTPCAAX1 /DB_XREF=g1:1777754 /UG-Hs.227777 protein tyrosine phosphatase type IVA, member 1 / EL=ab:U48296.1 qb:NM 003463.1 |
| 211372_s_at | soluble type II interleukin-1 receptor mRNA, complete cds. /FEA-mRNA /PROD-soluble type 188065 /UG-Hs.25333 interleukin 1 receptor, type II /FI-gb:U64094.1 |
| 204794_at | 5 P |
| 206177_s_at | gb:NM 000045.2 /DEE-Homo sapiens arginase, liver (ARG1), mRNA. /FEA-mRNA /GEN-ARG1 /PROD-arginase, type I /DB_XREF-gi:10947138 / UG-HS.289057 arginase, liver /FL-gb:NM 000045.2 gb:M14502.1 |
| 205239_at | io sapiens amphiregulin (schwannoma-derived growth factor) (AREG), mRNA. /FE vannoma-derived growth factor) /DB_XREF=gi:4502198 /UG-Hs.270833 amphiregulir cgb:NM 001657.1 |
| 207094_at | |
| 200776_s_at | 18328 /FEA-EST /DB XREF-g1:12781821 /DB_XREF-est:AL518328 /CLONE-CSODA009YK18 (3 p .gb:D13630.1 gb:NM 014670.1 |
| 215009_s_at | 192014.1 /DEE=Human clone =Hs 153527 Homo sapiens p |
| 211302_s_at | <pre>complete cds. /FEA=mRNA /FROD=phosphodiesterase /DB_XREF=g1:34712 (dunce (Drosophila)-homolog phosphodiesterase E4) /FL=gb:L20966.1</pre> |
| 210512_s_at | gb:AF022375.1 /DEF-Homo sapiens vascular endothelial growth factor mRNA, complete cds. /FEA-mRNA /PROD-vascular endothelial growth factor /DB %REF-gi:3719220 /UG-Hs.73793 vascular endothelial growth factor /FL-gb:M32977.1 gb:AF022375.1 gb:NM 003376.1 gb:AE021221.1 gb:AF091352.1 |
| 212577_at | AA868754 protein |
| 214866_at | ns mRNA for urokinase plasminogen activator receptor. /FEA=mRNA /PROD-urokinase 456192 /UG=Hs.179657 plasminogen activator, urokinase receptor |
| 219228_at | 'C2H2-11ke zinc finger protein (ZNF361), mRNA. /FEA-mRNA /GEN-ZNF361 /F *Hs.147644 zinc finger protein 331 /FL-qb:AF251515.2 qb:NM 018555.2 gb: |
| 201574_at | translation terminati tor 1 /DB_XREF=gi:4759 |
| 209498_at | X16354.1 /DEF-Human mRNA for transmembrane carcinoembryonic antigen BGPa (formerly TM1-CEP, preprotein /DB XREF-g1:37197 /UG-Hs.50964 carcinoembryonic antigen-related cell adhesion /FL-qb:J03858:1 |
| 207630_s_at | Dmo sapiens CAMP responsive element modul XREF-q1:4503038 /UG-Hs.155924 CAMP respo |
| 210873_x_at | sapiens phorbolin I mRNA, complete cds. /FEA=n apolipoprotein B mRNA editing protein) /FL=gb |
| 204419_x_at | sapiens hemoglobin, gamma G (HBG2), mRNA. /FEA-mRNA /GEN-HBG2 /PROD-hemoglobir, 1.283108 hemoglobin, gamma G /FL-gb:NM 000184.1 |
| 202988_s_at | of G-protein signa /UG=Hs.75256 regul |
| 209273_s_at | BG387555 /FEA-EST /DB XREF-91:13281001 GG4276 similar to CG8198 /FL-9b:AF2847 |
| 202693 s at | AN194730 /FEA-EST /DB XREF-gi:6473630 /DB XREF-est:xx43d11.x1 /CLONE=1MAGE:2696469 /UG-HS.90/5 17a (apoptos1s-inducing) /FI-gb:AB011420.1 gb:NM 004760.1 |
| 205270_s_at | gb:NN 005565.2 /DEF=Nomo sapiens lymphocyte cytosolic protein 2 (SH2 domain-containing Leukocyte protein of 76kD) (LCF2), mKNA. /FEA=mRNA /GEN=LCP2 /PROD=lymphocyte cytosolic protein 2 /DB XREE=gl:7382491 /UG=Hs.2488 lymphocyte cytosolic protein 2 (SH2 domain-containing leukocyte protein of 76kD) /FL=gb:NM 005565.2 gb:U20158.1 |

| Ν | 202284_s_at | tinase inhibitor LA (p2) |
|----------|-------------|---|
| | 219471_at | mo sapiens hypothetical protein FLJ21562 (FLJ21562), mRNA. /FEA=mRNA /GEI REF=g1:13376686 /UG=HS.208708 hypothetical protein FLJ21562 /FL=qb:NM 02: |
| | 208651_x_at | apiens CD24 sign s.286124 CD24 an |
| 101 | | 9b:AF015524.1 /DEF-Homo sapiens putative chemokine receptor (CRAM-A) mRNA, complete cds. /FEA-mRNA /GEN=CRAM-A / PROD-putative chemokine receptor /DB XREF-q1:3550066 /UG-Hs.302043 chemokine (C-C motif) receptor-11ke 2 /FL-ab.aF015524 1 |
| 102_ | 204285_s_at | 1857639 /FEA=EST /DB XREF=g1:5511255 /DB XREF-est:wk95g09.x1 /CLONE=IMAGE:2423200 / -acetate-induced profein 1 /FL=qb:NM 021127.1 |
| [0 | 221824_s_at | A770170 /FEA=EST /DB_XREF=q |
| <u> </u> | | 9b:NM 024668.1 /DEE-Homo sapiens hypothetical protein FLJ20288 (FLJ20288), mRNA. /FEA-mRNA /GEN-FLJ20288 /PROD-hypothetical protein FLJ20288 /FL-qb:BC004457.1 db:NM 024668.1 |
| 7 | | ISA238982 /PROD=VNN3 prot |
| <u>~</u> | | gb:NM 003059.1 /DEF=Homo sapiens solute carrier family 22 (organic cation transporter), member 4 (SLC22A4), mRNA. / FER=mRNA /GEN=SLC22A4 /PROD=solute carrier family 22 (organic cationtransporter), member 4 /DB XREF=g1:4507002 / UG-Hs.77239 solute carrier family 22 (organic cation transporter), member 4 /FL=qb:AB007448.1 db:NM 003059.1 |
| | 211445_x_at | sapiens FKSG17 (FKSG17) mRNA, complete cds. /FEA-mRNA /GEN-FKSG17 /PROD-FKSG ins FKSG17 (FKSG17) mRNA, complete cds /FL-gb:AF315951.1 |
| | 204286_s_at | gb:NM_021127.1 /DEF=Homo sapiens phorbol-12-myristate-13-acetate-induced protein 1 (PMAIPI), mRNA. /FEA=mRNA /GEN=PMAIPI / PROD=phorbol-12-myristate-13-acetate-induced protein1 /DB_XREF=gi:10863922 /UG-Hs.96 phorbol-12-myristate-13-acetate-induced protein 1 /FLa-ob:NN 021127.1 |
| 100 | | 9b:NH 014736.1 /DEF=Homo sapiens KIAA0101 gene product (KIAA0101), mRNA. /FEA=mRNA /GEN-KIAA0101 /FROD-KIAA0101 gene product / DB XREF=g1:7661905 /UG-Hs.81892 KIAA0101 gene product /FI-ap:D14657.1 ap:NM 014736.1 |
| īcī | 211560_s_at | sapiens clone FLB8929 PRO2399 5 outer mitochondrial membrane |
| | 209945_s_at | o sapiens, Similar to gl to glycogen synthase ki |
| | 217997_at | Consensus includes gb:AI795908 /FEA-EST /DB XREF=g1:5361371 /DB XREF=est:wh40a05.x1 /CLONE-IMAGE:2383184 /UG-Hs.82101 pleckstrin homology-like domain, family A, member 1 /FI-gb:NM 007350.1 |
| | 203757_s_at | mo sapiens, carcinoembryonic antigen-r complete cds. /FEA-mRNA /PROD=carci |
| | | (non-specific cross reacting antigen) /DB_XREF-gi:13477106 /UG-Hs.73848 carcinoembryonic antigen-related cell adhesion molecule 6 (non-specific cross reacting antigen) /FL-gb:BCO05008.1 qb:M18216.1 qb:M29541.1 qb:MN 002483.1 |
| N_ | 205269_at | Consensus includes gb:AI123251 /FEA=EST /DB XREF=g1:3539017 /DB XREF=est:qa47g03.x1 /CLONE=IMAGE:1689940 /UG=H8.2488 lymphocyte cytosolic protein 2 [SH2 domain-containing leukocyte protein of 76kD) /FIE-ch:NN 005565.2 ch:U20158 1 |
| 101 | 219049_at | no sapiens hypothetical protein FLJ11264 (FLJ11264), mRNA. /FEA-mRNA /G |
| 2 | 209396_s_at | glycoprotein mRNA, complete cds. 3-like 1 (cartilage glycoprotein- |
| 101 | 202637_s_at | II608725 /FEA-EST /DB XREE-gi:4617892 /DB XREF-est:tw? .ular adhesion molecule 1 (CD54), human rhinovirus rec |
| : | 205557_at | 9b:NM 001725.1 /DEF-Homo sapiens bactericidalpermeability-increasing protein (BPI), mRNA. /FEA-mRNA /GEN-BPI /PROD-bactericidalpermeability-increasing proteinprecursor /DB XREF-gi:4502446 /UG-Hs.89535 bactericidalpermeability-increasing protein /FL-qb:AF322588.1 qb:304739.1 qb:NN 001725.1 |
| | 207072_at | rotein (<u>IL18RAP</u>), mRMA. in /DB_XREF=gi:4504656 346.1 gb:NM 003853.1 |

| 202498_s_at (| BE550486 /FEA-EST /DB XREF-q1:9792178 /DB XREF-est:7a2 rier family 2 (facilitated glucose transporter), membe |
|---------------|---|
| 218739_at | CGI-58 protein (LOC51099), mRNA, /FEA=mRNA /GEN=LOC51099 /PROD=C CGI-58 protein /FL=gb:AF151816.1 gb:NM 016006.1 |
| at | ** |
| | |
| 200985_s_at | qb:NM 000611.1 /DEF=Homo sapiens CD59 antigen p18-20 (antigen teentilled by monoclonal antibooles 16.345, bulo, EJ30, E132 and G344) (CD59), mRNA. /FER=mRNA /GEN=CD59 /PROD=CD59 antigen p18-20 (antigen identified by monoclonal antibooles 16.345, EJ16, EJ30, EJ32 and G344) /DB XREF=g1:10835164 /UG=HS:119663 CD59 antigen p18-20 lantioen identified by monoclonal antibooles 16.345, EJ16, EJ30, EJ32 and G344) /FL=qp:NM 006611.1 qb:M34671.1 |
| 206026_s_at | mo sapiens tumor necrosis factor, alpha-induced protein 6 (TNFAIP6), mRNA. /FEA or necrosis factor, alpha-induced protein 6 /DB_XREF=gi:6005905 rosis factor, alpha-induced protein 6 /Fir-pb:NM 007115.1 |
| | o sapiens -kinase / |
| 202333_s_at | AA877765 /FEA=EST /DB XREF=g1:2886/30 /DB XREF=eET:nTUBIUS.SI /CLUNE=IMAGE:110103 /UG=RS.GII enzyme E2B (RAD6 homolog) /FI=gp:N74525.1 gb:NM 003337.1 |
| 201566_x_at | mRNA for Id-2H, complete cds. /FEA-mRNA /GEN-Id-2H /PROD=Id-2H /DB XREF- ng 2, dominant negative helix-loop-helix protein /FI-gb:M97796.1 gb:NM 0 |
| 44790_s_at | Cluster Incl. AI129310:qc48a05.x1 Homo sapiens CDNA, 3. end /clone=IMAGE-1712816 /clone_end=3 /gb=AI129310 /g1=359/824./ ug=Hs.234923 /len=811 |
| 213836_s_at | ANO52084 /FEA-EST /DB_XREF-gi:5914443 /DB_XREF-est:wy86f07.xl /CLONE-IMAGE:2555461 /UG- |
| 205027_s_at | D5204.1 /DEF-Homo sapiens mitogen-activated protein kinase kinase kinase 8 (MA togen-activated protein kinase kinase kinase8 /DB_XREF-gi:4885146 /UG-Hs.248 m 8./EL-qb:D14497.1 qb:NM 005204.1 |
| 207329_at | omo sapiens matrix metalloproteinase 8 (neutrophil collagenase) (MMP8), mRNA. /PROD=matrix metalloproteinase 8 préproprotein /DB_XREF-g1:4505220 etalloproteinase 8 (neutrophil collagenase) /FL-gb:J05556.1 gb:NM 002424.1 |
| | . A4576961 /FEA-EST /DB_XREF=g1:2354435 /DB.XREF-est:nm82d08.s1 /CLONE-IMAGE:1074/3 ike domain, family A, member 1 /FL-qb:NN 007350.1 |
| 208632 at | DB_XREF=est:AL578551 /CLONE=CSUDKUU1YGUI (3 |
| 206851_at | <pre>RNase A family, 3 (eosinophil cationic protein) (KNASE3), A family, 3 (eosinophilcationic protein) / DB XREF=g1:45065 osinophil cationic protein) / FE=gb:NM 002935. I gb:M28128.1</pre> |
| 203434_s_at | A1433463 /FEA=EST /DB_XREF=gi:4289355 /DB_XREF=est:ti65911.xl:/CLONE=INAGE:2136937 /UG=Hs.1298 (neutral endopeptidase, enkephalinase, CALLA, CD10) /FL=qb:J03779.1 gp:NM 007287.1 gb:NM 007288 |
| 216236_s_at | om cione Ukrzpowykie/1/2 partial cos. 75 lute carrier family 2 (facilitated glucos |
| 206342_x_at | MA. /FEA-mR Hunter syndr |
| 201329_s_at | /DEE-Homo sapiens v-ets avian erythroblastosis virus E26 oncogene homolog 2 (E7S2), mRNA. an erythroblastosis virus E26 oncogenehomolog 2 /DB_XREE=g1:4885220 /UG-Hs.33146 v-ets avia omolog 2 /E1—gb:J04102.1 gb:NM 005239.1 |
| 200731_s_at | 3 XREF=g1:6391485 /DB XREF=est:xf43a12.x1 /CLONE=IMMCE:2620/98 /Uu=HS.2 =gb:U48296:1 gb:NH 003463.1 |
| 212508_at | Consensus includes gb:AK024029.1 /DEE-Homo sapiens CDNA FLJ13967 fis, clone Y/9AALUU1402, weakly similar to homo sapiens paraneoplastic cancer-testis-brain antigen (MA4) mRNA. /FEA-mRNA /DB XREF-gi:10436287 /UG-Hs.24719 modulator of apoptosis 1 / |

| | FL=9b:AF305550.1 gb:NM_022151.1 | |
|-------------|--|---|
| 208868_s_at | Consensus includes gb:BF125756 /FEA-EST /DB XREF-g1:1096479 sapiens mRNA; cDNA DKF2p564N1272 (from clone DKF2p564N1272) | /DB_XREF=qi:10964796 /DB_XREF=est:601763146F1 /CLONE=IMAGE:4026010 /UG-Hs.282654 Homo |
| 204054 at | gb.NM_000314.1 /DEF-Homo saptens phosphatase and tensin homolog (mutated in multiple / FEA-mRNA /GEN-FTEN / FROD-phosphatase and tensin homolog (mutated inmultiple advance / UG-HS.10712 phosphatase and tensin homolog (mutated in multiple advanced cancers 1) / FL-ab: 092436.1 db: 0405051.1 db: U95180.1 db: NM_000314.1 | omo sapiens phosphatase and tensin homolog (mutated in multiple advanced cancers 1) (PTEN), mRNA. PROD-phosphatase and tensin homolog (mutated inmultiple advanced cancers 1) /DB_XREF=gi:4506248 ses and tensin homolog (mutated in multiple advanced cancers 1) 3051.1 ab:195180.1 ab:NN 000314.1 |
| 212262_at | Consensus includés gb:AA149639 /FEA=EST /DB XREF=qi: mouse qualing QKI (KH domain RNA binding protein) /F | consensus includes gb: AA149639 /FEA=EST /DB XREF=gi:1720440 /DB XREF=est:2139606.s1 /CLONE=INAGE:504298 /UG-Hs.15020 homolog of nouse quaking QKI (KH domain RNA binding protein) /FL-gb:AF142419.1 gb:AF142422.1 |
| 208785_s_at | Consensus includes gb:BE893893 /FEA-ESI /DB XREF-gi: microtubule-associated proteins IAIB light Chain 3 / | Consensus includes gb:BE893893 /FEA-EST /DB XREF-g1:10355716 /DB XREF-est:601436260F1 /CLONE-IMAGE:3921446 /UG-Hs.121849 microtubule-associated proteins lAIB 11ght chain 3 /FI-gb:AF303888.1 gb:NM 022818.2 gb:AF183417.1 |
| 218660_at | jb:NM 003494.1 /DEF=Homo sapiens dysferlin, limb girdle muscular dystrophy GEN=DYSF /PROD=dysferlin, DB XREF=g1:4503430 /UG=Hs.143897 dysferlin, limb FL=qb:AF075575.1 gb:NM 003494.1 | limb girdle muscular dystrophy 2B (autosomal recessive) (DYSF), mRNA. /FEA=mRNA / VG-Hs.143897 dysferlin, limb girdle muscular dystrophy 2B (autosomal recessive) / |
| 221920_s_at | Consensus includes gb:BE617761 /FEA-EST /DB_XREF-gi: //UG-Hs.300496 mitochondrial solute carrier | Consensus includes gb:BE617161 /FEA-EST /DB_XREE-gi:10038376 /DB_XREE-est:7£59h05.x1 /CLONE-IMAGE:3299001 /UG-Hs.300496 mitochondrial solute carrier |
| 211982_x_at | Consensus includes gb:AL546600 /FEA-EST /DB_XREF-g1: /UG-Hs.70500 KIAA0370 protein | Consensus includes gb:AL546600 /FEA-EST /DB_XREF-g1:12879872 /DB_XREF-est:AL546600 /CLONE-CS0DI029YA23 (3 prime) /uG-Hs 70500 KIAA0370 protein |
| 218647_s_at | gb:NM 024640.1 /DEF~Homo sapiens hypothetical protet PROD-hypothetical protein FLJ23476 /DB XREF~g1:13375 | gb:NM 024640.1 /DEF=Homo sapiens hypothetical protein FLJ23476 (FLJ23476), mRNA. /FEA=mRNA /GEN=FLJ23476 PROD-hypothetical protein FLJ23476 /DB XREF=gi:13375875 /UG-Hs.46736 hypothetical protein FLJ23476 /FL=gb:NM 024640.1 |
| 206571_s_at | gb:NM_004834.1 /DEF=Homo sapiens mitogen-activated protein kinase ki /FEA=mRNR /GEN=MAP4K4 /PROD=mitogen-activated protein kinase kinase /UG-Hs.3628 mitogen-activated protein kinase kinase kinase kinase | mo sapiens mitogen-activated protein kinase kinase kinase kinase 4 (MAP4K4), mRNA. /PROD-mitogen-activated protein kinase kinase kinasekinase 4 /DB XREF-qi:4758523 tivated protein kinase kinase kinase kinase 4 /FL-qb:AF096300.1 qb:NM 004834.1 |
| 218332_at | <pre>gb:NN 018476.1 /DEF=Homo sapiens uncharacterized hypothal. PROD=uncharacterized hypothalamus protein HBEX2 /DB XREF= FI=gb:AF220189.1 gb:NN 018476.1 gb:AF183416.1 gb:AF237783</pre> | amus protei gi:8923715 .1 |
| 207113_s_at | <pre>gb:NN 000594.1 /DEF=Homo saplens tumor necrosis factor (INF PROD=Tumor necrosis factor (cachectin) /DB_XREF=gi:10835154 FL=gb:NN 000594.1</pre> | or (INE superfamily, member 2) (TNF), mRNA. /FEA-mRNA /GEN-INF / 0835154 /UG-Hs.241570 tumor necrosis factor (TNE superfamily, member 2) / |
| 209388_at | <pre>gb:BC000927.1 /DEF-Homo sapiens, Similar to poly (A) /FEA-mRNA /PROD-Similar to poly (A) polymerase /DB X</pre> | Similar to poly (A) polymerase, clone NGC:5378, mRNA, complete cds. (A) polymerase /DB XREF-gi:12654216 /UG-Hs.49007 poly(A) polymerase alpha /FL-qb:BC000927.1 |
| 207610_s_at | <pre>gb:iM4 013447.1 /DEF=Homo saplens egf=like module con /FER=mRNA /GEN=EMR2 /PROD=egf=like module containin /UG=Hs.137354 egf=like module containing, mucin-like</pre> | gb:NM 013447.1 /DEF-Homo sapiens egf-like modula containing, mucin-like, hormone receptor-like sequence 2 (ENRZ), mRNA. /FER-mRNA /GEN-ENR2 /PROD-egf-like modula containing, mucin-like, hormonereceptor-like sequence 2 /DB XREF-gi:7305024 /UG-Hs.137354 egf-like module containing, mucin-like, hormone receptor-like sequence 2 /FI-qb:AF114491_1 qb:NM 013447.1 |
| 204258_at | <pre>gb:NN 001270.1 / DEF=Homo saplens chromodomain helica /PROD=chromodomain helicase DNA binding protein 1 / D binding protein 1 / FL-qb:AF006513.1 qb:NM 001270.1</pre> | jb:NM 001270.1 /DEF=Homo sapiens chromodomain helicase DNA binding protein 1 (CHDI), mRNA. FEA=mRNA /GEN=CHDI PROD=chromodomain helicase DNA binding protein 1 /DB_XREF=gi:4557446 /UG=Hs.22670 chromodomain helicase DNA binding protein 1 /FL=qb:AF006513.1 qb:NM 001270.1 |
| 200948_at | <pre>gb:NM 005439.1 /DEF=Homo sapiens myeloid leukemia factor 2 (MLF2), mRNA, /PROD=myeloid leukemia factor 2 /DB XREF=gi:4885486 /UG=Hs.79026 myeloid /FL=qb:BC000898.1 qb:BC002340.1 qb:\u057342.1 qb:AF070539.1 qb:NM 005439.1</pre> | . myeloid leukemia factor 2 (MLF2), mRNA. /FEA-mRNA /GEN-MLF2 /DB XREF-gi:4885486 /UG-Hs.79026 myeloid leukemia factor 2 qb:US7342.1 qb:AE070539.1 qb:NM 005439.1 |
| 211506_s_at | gb:AF043337.1 /DEF=Homo saplens interleukin 8 C-terminal variant (IL8) mRNA, complete cds. /GEN=IL8 /PROD=interleukin 8 C-terminal variant /DB XREF=gi:12641914 /UG=Hs.624 interleuki | mo sapiens interleukin 8 C-terminal variant (IL8) mRNA, complete cds. /FEA-mRNA eukin 8 C-terminal variant /DB XREF-gi:12641914 /UG-Hs.624 interleukin 8 /FL-gb:AF043337.1 |
| 217966_s_at | gb:NM 022083.1 /DEF=Homo sapiens niban procein (NIBAN), mRNA. /FEA=mRNA /GEN=NIBA /DB ÄREF=gi:11545796 /UG-Hs.48778 niban protein /FL-gb:AB050477.1 gb:NM 022083.1 | (NIBAN), mRNA. /FEA=mRNA /GEN=NIBAN /PROD=niban protein n /FL=gb:AB050477.1 gb:NM 022083.1 gb:AF288391.1 |
| 214151_s_at | Consensus includes gb:AU144243 /FEA-EST /DB XREF-gi:11005764 /DB_XREF-est:AU144243 /UG-Hs.247118 phosphatidylinositol glycan, class B | 11005764 /DB_XREF=est:AU144243 /CLONE=HEMBA1001328 |
| 211810_s_at | <pre>gb:DZ5284.1 /DEF=Human mRNA for alternative spliced product of /FEA=CDS /PROD=alternative spliced product of galactocerebrosidase /DB XREF=gi:457445 /UG=Hs.273 g</pre> | alternative spliced product of galactocerebrosidase, complete cds. ed /DB XREF-gi:457445 /UG-Hs.273 galactosylceramidase (Krabbe disease) /FL-gb:D25284.1 |
| 207890_s_at | | inase 25 (MMP25), transcript variant 2, mRNA. /FEA-mRNA rotein /DB_XREF-gi:13027808 022718.1 |

| 37028_at | Cluster Incl. U83981: Homo sapiens apoptosis associated protein (GADD34) mRNA, complete cds /cds=(222,2246) /gb=U83981 / n=3288617 /uo=Hs. 76556 /lon=2331 |
|-------------|---|
| 40446_at | cl. AL021366:cI |
| 219062_s_at | mo sapiens hypothetical protein FLJ20281 (FLJ20281), tein FLJ20281 /DB XREF-gi:8923259 /UG-Hs.18800 hypoth |
| 202158_s_at | KNA-binding procein z (CUGBEZ), mkNA. RREF-gi:5729815 /UG-Hs.211610 CUG triple jb:AF090694.1 gb:NM 006561.1 |
| 60084_at | itj6lell.xl Homo sapiens c 68 /len=554 |
| 210142_x_at | no sapiens flotillin mRNA, complete cds. /FEA-mRNA /PROD-flotillin /DB_XREF-gi: In 1 /FL-gb:AF117234.1 |
| 204490_s_at | i CDw44 antigen, complete cds. /FEA-mRNA /DB XREF-gi:180196 /UG-Hs.169610 CD44 antigen, complete cds. /FEA-mRNA /DB XREF-gi:180196 /UG-Hs.169610 Lgb:M24915. |
| 203949_at | gb:NM_000250.1 /DEF=Homo sapiens myeloperoxidase (NFO), nuclear gene encoding mitochondrial protein, mkWA. /FER=mRNA /GEN=NPO /PROD=myeloperoxidase /DB_XREF=g1:4557758 /UG=Hs.1817 myeloperoxidase /EL=gb:MJ9507.1 ob:JD2694.1 ob:NM:000250.1 |
| 219259_at | mmo sapiens hypothetical protein FLJ12287 similar to semaphorir 37 /PROD-hypothetical protein FLJ12287. similar tosemaphorins /I 31 protein FLJ12287 similar to semaphorins /FL-gb:NM 022367.1 |
| 201751_at | gb:NN:014876.1 /DEF-Homo sapiens KIRA0063 gene product (KIRA0063); mRNA. /FEA-mRNA /GEN-KIRA0063 /PRODE-KIRA0063 gene product /DB XREF-gi:7661887 /UG-Hs:3094 KIRA0063 gene product /FL-gb:D31884.1 gb:NM 014876.1 |
| 203153_at | /DEF=H =IFIT1 nterfer |
| 210724_at | eceptor EMR3 mRNA, comp XREF=gi:13183148 or EMR3 mRNA, complete |
| 202381_at | pb:NM 003816.1 /DEF-Homo sapiens a disintegrin and metalloproteinase domain 9 (melrrin gamma) (ADAM9), mRNA. /FEA-mANA /CEN-ADAM9 /FROD-a disintegrin and metalloproteiniase domain 9 perproprotein /DB XREF=gi:4501914 |
| 217824_at | ::UI-HF-BNO-aki-e- NF161502.1 gb:AF15 |
| 200984_s_at | Consensus includes gb:X16447.1 /DEF-Human mRNA for CD59, an LY-6-11He protein regulating complement membrane attack. / FEA-mRNA /PROD-precursor polypeptide (AA'-25 to 103) /DB_XREF-gi:29805 /UG-Hs.119663 CD59 antigen p18-20 (antigen identified by monoclonal antibodies 16.3A5, EJ16, EJ32, and G344) /FL-gb:NM 000611.1 gb:M34671.1 |
| 219999_at | in PRO2198 (PRO2198), mRNA. /FEA- 129 /UG-HS.116459 hypothetical px |
| 212644_s_at | Clone HEP04178, highly simi |
| 219622_at | omo sapiens hypothetical protein FLJ20429 (FLJ20429), mRNA. /FEA=mRNA / otein FLJ20429 /DB XREE-gi:8923400 /UG-Hs.179791 hypothetical protein |
| 219190_s_at | gb:NN 017629.1 / DEF-Homo sapiens hypothetical protein FL/320033 (FL/320033), mRNA. / FEA-mRNA. / GEN-FL/320033 FL/320033 / FL/32003 |
| 208707_at | :BE552334 /FDA=EST /DB XREF=g1:9/94026 /DB XREF=eST:Ny tic translation initiation factor 5 /FL=gb:ALO80102.1 |
| 218506_x_at | omo sapiens uncharacterized bone marrow piccein Emv43 (Bru43), aracterized bone marrow protein BM045 (DB XREF-gi:8922103 erized bone marrow protein BM045 (FL-gb:AF21752):1 gb:NM 01845 |
| 207545_s_at | pb:NM_003744.1 /DEF-Homo sapiens numb (Drosophila) homolog (NUMB), mRNM. /FEA-mRNA /GEN-NUMB /FKOD-numb (Drosophila) nomolog / DB XREF-gi:4505478 /UG-Hs.78890 numb (Drosophila) homolog /FL-gb:NM 003744.1 gb:L40393.1 |

| 212014_x_at | Consensus includes 9b:AI493245 /FEA=EST /DB_XREF=gi:4394248 /DB_XREF=est:ti30d08.x1 /CLONE=INAGE:2131983 //UG-Hs.169610 CD44 antigen (homing function and Indian blood group system) |
|-------------|--|
| 209039_x_at | gb:AF001434.1 /DEF-Human Hpast (HPAST) mRNA, complete cds. /FEA-mRNA /GEN-HPAST /PROD-Hpast /DB_XREF-g1:2529706 //UG-Hs.155119 EH domain containing 1 /FL-gb:AF001434.1 |
| 208650_s_at | Consensus includes gb:BG327863 /FEA-EST /DB_XREE-g1:13134301 /DB_XREE-est:602426876F1 /CLONE=1NAGE:4564675 // OFFEA OFFE |
| 209806_at | omo sapiens, H2B histone family, member A, clone NGC:5132, mRNA, complete cds. /FEA-mR nily, member A /DB XREF-g1:12654150 /UG-Hs.247817 H2B histone family. member A /FI-ch.F |
| 212573_at | /DEF-Homo sapiens clone 24951 mRNA sequence. /FEA-mRNA /DB_XREF-gi: |
| 203845_at | Consensus includes gb:AV727449 /FEA-EST /DB XREF=g1:10836870 /DB XREF=est:AV727449 /CLONE-HTCAYG01 /UG-Hs.199061 p300CBP-associated factor /FL-gb:U57317.2 gb:NM 003884.2 |
| 215806_x_at | Consensus includes gb:M13231.1 /DEF-Human T-cell receptor aberrantly rearranged gamma-chain mRNA from cell line HPB-MLT. /FEA-mRNA /DB XREF-gi:339168 /UG-Hs.274509 T cell receptor gamma constant 2 |
| 202195_s_at | Homo sapiens CGI-100 protein (LOC50999), mRNA. /FEA-mRNA /GEN-LOC /UG-Hs.296155 CGI-100 protein /FL-gb.AF151858.1 qb:NN 016040.1 |
| 201912_s_at | gb:NM 002094.1 /DEE-Homo sapiens G1 to S phase transition 1 (GSPT1), mRNA. /FEA-mRNA /GEN-GSP71 // PROD-G1 to S phase transition 1 /DB XREF-g1:4504166 /UG-Hs.2707 G1 to S phase transition 1 /FE-dp:NM 002094.1 |
| 201712_s_at | 101 5 |
| 212864_at | o:Y16521.1 /DEF=Homo sapiens mRNA for CDS2 protein. /FFA=mRNA /GEN=cds2 DB XREF=q1:4186022 /UG=Hs.24812 CDP-diacylq1ycerol synthase (phosohatidate cyridyly |
| 203504_s_at | "Homo sapiens ATP-binding cassette, sub-family A (ABC1), member 1 (ABCA1), m Al /PROD=ATP-binding cassette, sub-family A member 1 /DB XREF=q1:591567 |
| | ding cassette, sub-family A (ABC1), member 1 /FL-gb:AFI |
| 212457_at | Consensus includes gb:AL161985.1 /DEF=Homo sapiens mRNA; cDNA DKF2p761J1810 (from clone DKF2p761J1810). /FEA=mRNA /DB XREF=gi:7328121 /UG=Hs.274184 transcription factor binding to IGHM enhancer 3 |
| 214683_s_at | b:AI251890 kinasel |
| 44783_s_at | Cluster Incl. R61374:yh15e02.s1 Homo sapiens cDNA, 3 end /clone=IMAGE-37665 /clone_end-3 /gb=R61374 /gi=832069 /ug-Hs.234434 /len=434 |
| 202252_at | gb:NM 002870.1 /DEF-Homo sapiens RAB13, member RAS oncogene family (RAB13), mRNA. /FEA-mRNA /GEN-RAB13 /PROD-RAB13, member RAS oncogene family /DE XREF-gi:4506362 /UG-Hs.151536 RAB13, member RAS oncogene family /FL-gb:BC000799.1 |
| 206025_s_at | Consensus includes gb:AN188198 /FEA-EST /DB XREF-g1:6462634 /DB XREF-est:xj93f03.xl /CLONE-IMAGE:2664797 /UG-Hs.29352 tumor necrosis factor, alpha-Induced protein 6 /FL-qb:NM 007115,1 |
| 203936_s_at | gb:NM_004994.1 /DEE-Homo sapiens matrix metalloproteinase 9 (gelatinase B, 92kD gelatinase, 92kD type IV collagenase) [MMMP9], mRNA. /FEA-mRNA /GEN-NMP9 /PROD-matrix metalloproteinase 9 preproprotein /DB XREF-g1:4826835 /UG-Hs.151738 matrix metalloproteinase 9 (gelatinase B, 92kD gelatinase, 92kD type IV collagenase) /Fr-gh. Tochon 1 ch. NA OAGGA 1 |
| 221523_s_at | 0:AL138717 /DEF |
| • • | of the gene for a yeast uniquitin conjugating enzyme UBCC nomotog, the gene for a possible GTP binding protein, La MACA (nascent-polypeptide-associated complex alpha polype/FEA-mRNA 2 /DB_XREF=g1:8894207 /UG-Hs.238679 Rag D protein /FE-agb:RM 021244.1 qb:RZ72036.1 qb:BC003988.1 |
| 215785_s_at | Consensus includes gb:AL161999.1 /DEE=Homo sapiens mRNA; cDNA DKFZp761H087 (from clone DKFZp761H087); partial cds. /FEA=mRNA /GEN=DKFZp761H087 /PROD=hypothetical protein /DB XREF=q1:7328000 /UG-Hs. 558501 p53 inducible protein |
| 217822_at | (LOC51729), mRNA. /FEA=mRNA. NPWBP /FL=gb:BC001621.1 qb:A |
| 202206_at | o:AN450363 /FEA-EST /DB XREF-g1:6991139 /DB XREF-est:UI-H-BI3-akn-d-02-0-UI.sI /CLONE=INAGE:2 oosylation factor-like 7 /FL-gb:BC001051.1 gb:AB016811.1 gb:NM 005737.2 |
| 202439_s_at | gb:NM 000202.2 /DEE-Homo saplens iduronate 2-sulfatase (Hunter syndrome) (IDS), transcript variant 1, mRNA. /FEA-mRNA /GEN-IDS /PROD=iduronate-2-sulfatase isoform a precursor /DB XREF-gi:5360215 /VG=Hs.172458 iduronate 2-sulfatase (Hunter syndrome) /FL-qb:N58342.1 qb:NM 000202.2 |
| | |

| 213056_at | /FEA-EST /DB_XREF-gi:11006540 /DB_XREF-est:AU145019 /CLONE-HENBAld |
|---------------|---|
| | Consensus includes gb:AK024212.1 /DEF-Homo sapiens CDNA FLJ14150 fis, clone MAMMA1003026, highly similar to Homo sapiens HSPC057 mRNA. /FEA-mRNA /DB XREF-gi:10435534 /UG-Hs.18825 XFAA0849 protein |
| | 6 |
| 3t | /FEA=EST /DB_XREF=g1:5811/56 /UB_XREF=est: alpha |
| 212219_at | onsensus includes gb:D38521.1 /DEE=Human mRNA for KIAA0077 gene, partial cds. /FEA=mRNA /GEN=KIAA0077 /DB XREF-gi:559329 /UG-Hs.112396 KIAA0077 protein |
| 214152_at | 4243 |
| 202948_at | /FEA-mRNA /GEN-ILIRI /FROD-Ir /FL-gb:M27492.1 gb:NM 000877 |
| 207978_s_at | 95.NM 006981.1 /DEF=Homo, sapiens nuclear receptor subfamily 4, group A, member 3 (NR4A3), mRNA. /FEA-mKNA. /GEM=NR4A3 /PROD=nuclear.receptor subfamily 4, group A, member 3 /DB_XREF=g1:11276070 /UG=Hs.80561 nuclear receptor subfamily 4, |
| 209304_x_at | gb:AF087853.1 /DEF-Homo sapiens growth arrest and DNA damage inducible protein beta (GADD45B) mRNA, complete cds. GP:AF087853.1 /DEF-HOMO sapiens growth arrest and DNA damage inducible proteinbeta /DB XREF-gi:12061050 INC-HS. 110771 growth arrest and DNA-damage-inducible, beta /FL-gb:AF087853.1 gb:AF078077.1. |
| 214783_s_at | BG177920 /FERWEST / DB_XREF=gi:12684623 / DB_XREF=est:602327839F1 /CLONE=IMAGE:44 A11 |
| 46270_at | |
| 221760_at | BG287153 /FEA-EST /DB_XREF-gi:13040709 /DB_XREF- se, alpha, class 1A, member 1 |
| 210118_s_at | 2 I |
| \$204781_s_at | gb:NM 000043.1 /DEF-Homo sapiens tumor necrosis factor receptor superfamily, member 6 (INFRSF6), mRNA. /FEA-mRNA /GEN-TNFRSF6 /PROD-apoptosis (APO-1) antigen 1 /DB_XREF-gi:4507582 /UG-Hs.82359 tumor necrosis factor receptor superfamily, member 6 /FI-ch:M67454.1 db:NM 000043.1 |
| 207677_s_at | gb.NN 013416.1 (DEF-Homo sapiens neutrophil cytosolic factor 4 (40kD) (NCE4), transcript variant 2, mRNA. /FEA-mRNA /GEN-NCF4 /PROD-neutrophil cytosolic factor 4 (40kD), isoform 2 /DB XREF-gi:7382492 /UG-Hs.196352 neutrophil cytosolic factor 4 (40kD) /FL-gb:BC002798.1 gb:AB025219.1 gb:NN 013416.1 |
| 206302_s_at | gb:NM 019094.1 /DEF-Homo sapiens nudix (nucleoside diphosphate linked moiety X)-type motif 4 (NUDT4), mRNA. /FER-mRNA /GEN-NUDT4 /FROD-nudix (nucleoside diphosphate linked moietyX)-type motif 4 /DB XREF-gi:10800135/UG=Hs.92381 nudix (nucleoside diphosphate linked moiety X)-type motif 4 /FL-gb:NM 019094.1 gb:AF191649.1 gb:AF191650.1 gb:AF191653.1 |
| 219460_s_at | gb:NM 017849.1 /DEF-Homo sapiens hypothetical protein FLJ2050/ (FLJ2050/), mkNA. /EEA-mkNA /GEN-FLJ2050/ /FNUD-Nypothetical protein FLJ2050/ /FL-gb:NM 017849.1 |
| 212225_at | |
| 204479_at | omo sapiens 'DB XREF=q1 |
| 216457_s_at | <pre>1 /DEF=Homo sapiens cDNA: FL. 1, subunit 1, 120kD</pre> |
| 212676_at | 93356 /FEA=EST /DB_XREF CDNA: FLJ23005 fis, clo |
| 203675_at | obindin 2 (NUCB2), mRNA. / 642.1 gb:AF052643.1 gb:AF0 |
| 221156_x_at | qb:NM 004748.1 /DEF=Homo sapiens cell cycle progression 8 protein (CPR8), mkNA. /FEA=mRNA. /UEN=CR8 /FRUJ=CELL CYCLE progression 8 protein /DE=gb:RFS /FRUJ=GB / DE:NM 004748.1 progression 8 protein /DE=gb:RFS / DE:NM 004748.1 |
| 207791_s_at | gb:NM_004161.1 / DEF=Homo sapiens KABI, member KAS oncogene idmily (rABI), mnka: / rea-mnka / Och-KABI / 1.100-1402. |

| | member RAS oncogene family /DB_XREF=g1:4758987 /UG=Hs.3642 RAB1, member RAS oncogene family /FL=gb:NM_004161.1 gb:M28209.1 |
|-------------|---|
| 200692_s_at | gb:NM 004134.1 /DEF=Homo sapiens heat shock 70kD protein 9B (mortalin-2) (HSPA9B), mRNA. /FEA=mRNA /GEN-HSPA9B /PROD=heat shock 70kD protein 9B (mortalin-2) /DB_XREF=gi:4758569 /UG=Hs.3069 heat shock 70kD protein 9B (mortalin-2) /FL=gb:BC000478.1 gb:L15189.1 gb:NM 004134.1 |
| 218936_s_at | |
| 213931_at | 10.5 |
| 218247_s_at | o sapiens hypotheti Hs.12830 hypothetic |
| | .F007132.1 /DEF-Homo sapiens clone 23551 mRNA sequence. |
| 208815_x_at | sapiens mRNA for hea 1:4579908 /UG-Hs.90093 |
| 218177_at | A293502 /FEA-EST /DB XREF-g1:1941036 /DB XREF-est AF281064.1 qb:NM 020412.1 |
| 200648_s_at | gb:NW_002065.1 /DEF-Homo sapiens giutamate-anmonia ligase (giutamine synthase) (GLUL), mRNA. /FEA-mRNA /GEN-GLUL / PROD-giutamate-ammonia ligase (glutamine synthase) /DB XREF-gi:4504026 /UG-Hs.170171 Glutamate-ammonia ligase (glutamine synthase) /FI-ch.NW_00068 |
| 202843_at | thelial diffe |
| 214578 s at | 7.1 gb:AL080081.1 gb:AB026908.1 gb:NM 012328.1 |
| 20,014 | |
| 217473_x_at | s includes gb:AF229163 / cds, alternatively spli |
| | on-coupled divalent metal ion t |
| 207624_s_at | gb:NM 000328.1 /DEF-Homo sapiens retinitis pigmentosa GTPase regulator (RPGR), mRNA. /FEA-mRNA /GEN-RPGR /PROD-retinitis pigmentosa GTPase regulator /DB XREF-gi:4506580 /UG-H8.153614 retinitis pigmentosa GTPase remulator / |
| 202778_s_at | = inc finner |
| 200653 s at | /UG=Hs.109526 zi |
| במחחם בשרי | caimodulin mkNA, comple lase kinase, delta) /FI |
| 218627_at | gb:NM 018370.1 /DEF-Homo sapiens hypothetical protein FLJ11259 (FLJ11259), mRNR. /FEA-mRNA /GEN-FLJ11259 /PROD-hypothetical protein FLJ11259 /PB XREF-g1:8922957 /UG-Hs.184465 hypothetical protein FLJ11259 /FL-ch:NM 018370.1 |
| 202059_s_at | ilpha 5) (KPNA1), mRNA. /FEA in alpha 1 (importin alpha 5 |
| 218132_s_at | 9b:NM 024075.1 /DEF-Homo sapiens LENGS protein (LENGS), mRNA. /FEA-mRNA /GEN-LENGS /PROD-LENGS protein / DB XREF-gi:13129061 /UG-Hs.15580 LENGS protein /FL-db:BC000944 2 ob:NM 024075,1 |
| 203265_s_at | A810268 /FEA=EST /DB XRE ivated protein kinase ki |
| 209694_at | 6-pyruvoyltetrahydropterin synthase (PTS) mRNA, complete cds. ropterin synthase /DB_XREF=gi:306438 /UG=Hs.366 6-pyruvoyltetr 00317.1 ab:117400.1 |
| 217591_at | gb:BF725121 /FEA=EST |
| 222309_at | Consensus includes gb:AW972292 /FEA=EST /DB_XREF=gi:8162138 /DB_XREF=est:ESŢ384381 /UG=Hs.292998 ESTs |

| 201627_s_at | gb:NN 005542.1 /DEE-Homo sapiens insulin induced gene 1 (INSIGI), mRNA. /FEA-mRNA /GEN-INSIG1 / PROD-Insulin induced gene 1 /DB XREF-gi:5031800 /UG-Hs.56205 insulin induced gene 1 /FL-gb:NN 005542.1 |
|-------------|--|
| 217249_x_at | 14544 /DEF-Homo sapiens BAC clone CTB-162B4 from 4 /FEA- BAC clone CTB-162B4 from 4 |
| 220496_at | FEA-mRNA /GEN-LOC5126 -like receptor-2 /FL- |
| 201096_s_at | ludas gb:AL537042 /FEA=EST /DB XREF=g1:12800535 /DB XREF=est:AL537042 /CLONE=CSODF017YF17 (5 LDP-ribosylation factor 4 /FL=gb:BC003364.1 gb:N35341.1 gb:NM 001660.2 |
| 201580_s_at | Consensus includes gb:AL54094 /FEA-EST /DB_XREF=gi:12876573 /DB_XREF=est:AL544094 /CLONE=CSODI004YG20 (3 prime) / UG-Hs.169358 hypothetical protein /FL-gb:NM 021156.1 |
| 201109_s_at | 12 /UG-Hs. |
| 203080_s_at | sapiens bromodomain adjacent to zinc finger domain, 2B (BAZZB), mKNA. /FEA=mKN ; to zinc finger domain, 2B /DB_XREF=gi:7304922 /UG=HS.8383 bromodomain adjacent 55.1 gb:NN 013450.1 |
| 202558_s_at | gb:NN 006948.1 /DEF-Homo sapiens stress 70 protein chaperone, microsome-associated, 60kD (STCH), mRNA. / FEA-mRNA. / GEN-STCH /PROD-stress 70 protein chaperone, microsome-associated, 60kD /DB_XREF-g1:5902125-/UG-Hs.288799 stress 70 protein chaperone, microsome-associated, 60kD /FL-gb:U04735.1 gb:NN 006948.1 |
| 201349_at | gb:NN 004252.1 /DEF-Homo sapiens solute carrier family 9 (sodiumhydrogen exchanger), isoform 3 requiatory factor 1 |
| • | er family 9 (sodium)ydrogen exchanger), isoform 3 regul jb:AF015926.1 gb:NN 004252.1 |
| 220603_s_at | /FER=mRNA /GEN=FLJ11175 / protein FLJ11175 /FL=gb:NM 01 |
| 35820_at | for GM2 activator protein /cds=UNKNOWN /gb=X62078 /gi=313158 /ug=Hs. |
| 201731_s_at | <pre>gb:NM 003292.1 /DEF=Homo sapiens translocated promoter region (to activated MET oncogene) (TPR), mRNA. /FEA=mRNA./ GEN=TPR /PROD=translocated promoter region (to activated METoncogene) /DB_XREE=gi:4507658 /UG=Hs.169750 translocated hypometer region (to activated MET oncogene) /FL=qb:NM 003292.1</pre> |
| .202414_at | mo sapiens excision repair cross-complementing rodent repair deficiency, complementat, complementat, complementation group G (Cockayne syndrome)) (ERCCS), mRNA. /FEA-mRNA /GEN-ERCCS / |
| | protein /DB XREF-gi:4503600 /UG-Hs.48576 excision repair cross-complementing rodent rep ation group 5 (xeroderma pigmentosum, complementation group G (Cockayne syndrome)) / 46.1 db:NM 000123.1 |
| 210422_x_at | gb:D50402.1./DEF-Human mRNA for NRAMP1, complete cds. /FEA-mRNA /PROD-Nramp /DB XREF-g1:1000996 /UG-Hs.182611 solute carrier family 11 (proton-coupled divalent metal ion transporters), member 1 /FI-gb:D50402.1 gb:D50403.1 |
| 58900 at | gb:NM 000578:1 gb:L32185.1 |
| | 46 /len=454 |
| 219890_at | arbonydrate-recognition domain, rectin, subeliamily m dependent, carbohydrate-recognition domain) lectin, (calcium dependent, carbohydrate-recognition domain |
| | qb:NM 013252.1 gb:AF139768.1 |
| 208112_x_at | mo sapiens EH domain containing 1 (EHD1), mRNA. /FEA-mRNA. -Hs.155119 EH domain containing 1 /FL-gb:AF099011.1 gb:NM (|
| 218655_s_at | mo sapiens hypothetical protein FLJ20291 (FLJ20291), mRNA. ein FLJ20291 /DB XREF=gl:8923270 /UG=Hs.8928 hypothetical p |
| 214230_at | R37664 /FEA-EST /DB XREF-gi:795120 /DE sion cycle 42 (GTP-binding protein, 29 |
| 201898_s_at | Consensus includes gb:AII26625 /FEA=EST /DB_XREF=gi:3595139 /DB_XREF=est:gdb3a12.X1 /LLUNt=1040621 / UG=Hs.80612 ubiquitin-conjugating enzyme E2A (RAD6 homolog) /EL=gb:M74524.1 gb:NM 003336.1 |

| 210773_s_at | gb:UB1501.1 /DEF-Human lipozin A4 receptor mRNA, complete cds. /FEA-mRNA /PROD-lipoxin A4 receptor /DB XREF-gi:1916075 / Uncaris, 99855 formyl peptide receptor-like 1 /FL-qb:M76672.1 qb:M84562.1 qb:M88107.1 qb:U81501.1 qb:AF054013.1 |
|-------------|--|
| | gb:NM 001462.1 |
| 32069_at | Cluster Incl. AE014515: Homo sapiens mRNA for KIAA0615 protein, complete cds /cds=(237,2927) /gb-AE014515 /gi=3327043 / ug-Hs.155972 /Len=3319 |
| 121_at | K69699 /FEATURE / DEFINITION=HSPAX8A H.sapiens Pax8 mRNA |
| 202053_s_at | gb:L47162.1 /DEF-Human fatty aldehyde dehydrogenase (FALDH) mRNA, complete cds. /FEA-mRNA /GEN-FALDH /PROD-fatty aldehyde dehydrogenase /DB_XREF-gi:1082035 /UG-Hs.159608 aldehyde dehydrogenase 3 family, member A2 /FL-gb:L47162.1 db:U46689.1 db:NN 000382.1 |
| 212335_at | NAVI67793 /FEA-EST /DB XREF-g1:6399401 /DB_XREF-est:xg56d07.xl /CLONE-IM NRNA sequence, alternatively spliced |
| 209813_x_at | T-cell receptor gamma chain VJCI-CII-CIII region mRNA, complete cds. Hs.112259 T cell receptor gamma locus /FL=gb:N16768.1 gb:RF151103.1 |
| 204747_at | gb:NN 001549.1 /DEF-Homo sapiens interferon-induced protein with tetratricopeptide repeats 4 (IFIT4), mRNA. /FEA-mRNA / GEN-IFIT4 /PROD-interferon-induced protein withtetratricopeptide repeats 4 /DB XREF-gi:4504586 /UG-Hs.181874 interferon-induced protein with tetratricopeptide repeats 4 /FL-gb:BC001383.1 gb:BC004977.1 gb:U52513.1 gb:AF026939.1 |
| 4" 202010 | pb:AF083470.1 pb:NM 001549.1 |
| 717773. ac | Consensus includes guinistatudes (remeda) (be anti-gilloutido) (be anti-catinoutido) (beneda) |
| 218880_at | |
| ررا | ins lipin 2 (LPIN2), mRUA. /FEA-mRNA /GEN-LPIN2 /FROD-lipin 2 /DB_XREF-gi 37436.1 gb:NM 014646.1 |
| 58780_s_at | Cluster Incl. R42449:yg02a07.s1 Homo sapiens cDNA, 3 end /clone=IMAGE-30831 /clone_end=3 /gb=R42449 /gi=817213 / ug=Hs.235831 /len=431 |
| 205281_s_at | 1 /DEF=Homo sapiens phosphatidylinositol glycan, class A (paroxysmal noc riant 1, mRNA. /FEA-mRNA /GEN-PIGA /PROD-phosphatidylinositol glycan, c |
| | /UG-Hs.51 phosphatidylinositol glycan, class A (paroxysmal nocturnal hemoglobinuria) /FL-gb:NM 002641.1 gb:011466.1 |
| 218251_at | gb.NM 021242.1 /DEF=Homo sapiens hypothetical protein STRAIT11499 (STRAIT11499), mRNA. /FEA=mRNA /GEN=STRAIT11499 / PROD=hypothetical protein STRAIT11499 /DB XREF=g1:10864050 /UG=Hs.236556 hypothetical protein STRAIT11499 /FL=gb:NM 021242.1 |
| 212481_s_at | |
| 213292_s_at | Consensus includes gb:AA908770 /FEA=EST /DB_XREF=gi:3048175 /DB_XREF=est:ol06a11.s1 /CLONE=IMAGE:1522652 / UG-Hs.283881 KIAA0713 protein |
| 221497_x_at | 95:BC005369.1 /DEF-Homo sapiens, chromosome 1 open reading frame 12, clone MGC:12484, mRNA, complete cds. / FEA-mRNA /PROD-chromosome 1 open reading frame 12 /DB XREF=gi:13529208 /UG-Ks.6523 chromosome 1 open reading frame 12 / Frachiarysovas 1 obiascy17176 1 obias 02051.1 obiasC005369:1 |
| 212185_x_at | Consensus includes gb:NM 005953.1 /DEF=Homo sapiens metallothionein 2A (MT2A), mRNA. /FEA=CDS /GEN=MT2A / PROD=metallothionein 2A /DB XREF=qi:5174763 /UG=Hs.118786 metallothionein 2A /FE-qb:NM 005953.1 |
| 203961_at | ALIS7398 /DEF-Human DNA sequence from clone RPII-56H7 o |
| · | 3 1 |
| 210986_s_at | gb:224727.1 /DEE-H.sapiens tropomyosin isoform mRNA, complete CDS. /FEA-mRNA /PROD-tropomyosin isoform /DB_XREF-g1:854188 / UG-Hs.77899 tropomyosin 1 (alpha) /FL-gb:224727.1 |
| 41386_i_at | :Human mRNA for KIAA0346 gene, partial |
| 201874_at | BE978611 /FEA-EST /DB XREF-g1:12345826 /DB XREF-est:602149061F1 /CLONE-I al protein FLJ21047 /FL-gb:NM 024569.1 |
| 211781_x_at | o sapiens, clone L≖gb:BC006164.1 |
| 202459_s_at | Consensus includes gb:U55968 /FEA=EST /DB_XREF=g1:1334524 /DB_XREF=est:HSU55968 /CLONE=26508 /UG=Hs.16b318 lipin 2 / FL=gb:D87436.1 gb:NM 014646:1 |

Table 4: Genes from Cluster Analysis 4

| 2 6 6 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 | |
|---|---|
| Arrymetrix internal designation | Description of the sequence in the GeneBank data base |
| 221798_x_at | Consensus includes gb:AI183766 /FEA=EST /DB_XREF=gi:3734404 /DB_XREF=est:qe17g06.x1 /CLONE=IMAGE:1739290 /UG=Hs.182426 ribosomal protein S2 |
| 21 <u>8231_</u> at | gb:NM 017567.1 /DEF=Homo sapiens N-Acetylglucosamine kinase (HSA242910), mRNA. /FEA=mRNA /GEN=HSA242910 /PROD=N- Acetylglucosamine kinase /DB XREF=gi:8923736 /UG=Hs.7036 N-Acetylglucosamine kinase /FL=gb:BC001029.1 gb:BC005371.1 gb:NM 017567.1 |
| 201009_s_at | Consensus includes gb:AI439556 /FEA=EST /DB_XREF=gi:4305149 /DB_XREF=est:tc90c12.x1 /CLONE=IMAGE:2073430 /UG=Hs.179526 upregulated by 1,25-dihydroxyvitamin D-3 /FL=gb:NM 006472.1 gb:S73591.1 |
| 212998_x_at | Consensus includes gb:AIS83173 /FEA=EST /DB_XREF=gi:4569070 /DB_XREF=est:tq64e04.x1 /CLONE=IMAGE:2213598 /UG=Hs.73931 major histocompatibility complex, class II, DQ beta 1 |
| 201464_x_at | |
| 214687_x_at | |
| at | gb:NM 014020.1 /DEF-Homo sapiens LR8 protein (LR8), mRNA. /FEA-mRNA /GEN-LR8 /PROD-LR8 protein /DB_XREF-gi:7662497 /UG-H5.190161 LR8 protein /FL-gb:AF115384.1 gb:NM 014020.1 |
| | gb:NM 001765.1 /DEF-Homo sapiens CD1C antigen, c polypeptide (CD1C), mRNA. /FEA=mRNA /GEN-CD1C /PROD-CD1C antigen, c po- lypeptide / DB XREF-gi:4502646 /UG-Hs.1311 CD1C antigen, c polypeptide /FL-gb:M28827.1 gb:NM 001765.1 |
| 202658_at | .1 /DEF=Homo sapiens peroxisomal biogenesis factor 11B (PEX11B), mRNA. /FEA=mRNA /GEN=PEX11B /PROD=peroxisomal biogenesis factor 11B /DB XREF=gi:4505718 /UG=Hs.83023 peroxisomal biogenesis factor 11B /FL=gb:AF093670.1 gb:AB018080.1 gb:NM 003846.1 |
| 200991_s_at | gb:NM 014748.1 /DEF-Homo sapiens KIAA0064 gene product (KIAA0064), mRNA. /FEA-mRNA /GEN-KIAA0064 /PROD-KIAA0064 gene product / OB XREF-gi:7661889 /UG-Hs.278569 sorting nexin 17 /FL-gb:BC002524.1 gb:BC002610.1 gb:D31764.1 gb:NM 014748.1 |
| 209007_s_at | gb:AF267856.1 /DEF=Homo sapiens HT033 mRNA, complete cds. /FEA=mRNA /PROD=HT033 /DB_XREF=gi:12006038 /UG=Hs.8084 hypo- thetical protein dJ465N24.2.1 /FL-gb:AF247168.1 gb:AF267856.1 |
| | Consensus includes gb:BF683426 /FEA=EST /DB_XREF=gi:11968834 /DB_XREF=est:602139603F1 /CLONE=IMAGE:4300777 /UG=Hs.183698 ribosomal protein L29 |
| | . H |
| 208929_x_at | gb:BC004954.1 /DEF=Homo sapiens, clone MGC:10897, mRNA, complete cds. /FEA=mRNA /PROD=Unknown (protein for MGC:10897) /DB XREF=gi:13436331 /UG-Hs.180842 ribosomal protein L13 /FL=gb:BC000851.1 gb:BC004954.1 gb:NM 000977.1 |

| 206170_at | |
|-------------|---|
| | /rkuu=adrenergic, beta-2-, receptor, surface /DB XREF=qi:13162366 /UG=Hs.2551 adrenergic, beta-2 receptor, surface /Fi=ah·NM 000024 2 ah·M15168 1 |
| 212582_at | includes gb: ALO49923.1 / DEF-Homo sapiens mRNA; cDNA DKF2p564E2282 (from clone DKF2p564E2282). /FERAgi: 4884169 /UG-Hs.109694 KIAA1451 protein |
| 208630_at | Consensus includes gb:AI972144 /FEA=EST /DB_XREF=gi:5768970 /DB_XREF=est:wr63b03.x1 /CLOWE=IMAGE:2492333 /UG=Rs.75860 hydroxyacyl-Coenzyme A dehydrogenase3-ketoacyl-Coenzyme A thiolaseenoyl-Coenzyme A hydratase /trifingtional profess and all the continuity of the |
| 214058_at | arpus subunit /in-go.na-vectori; guinting- 720 /DEF-Human L-myc protein gene, complete cd mwellocytomatchesis viral oncodene homolog 1 1 |
| 201991_s_at | 11130401 / FEA=EST / DB_XREF=g1:11130401 / DB_XREF=est:7q30g05.x |
| 212179_at | 01 /FEA=EST /DB_XREF= cotein |
| 210949_s_at | gb:BC000533.1 /DEF=Homo sapiens, Similar to eukaryotic translation initiation factor 3, subunit 8 (110kD), clone |
| | proc.093, mRNA, complete cds. /FEA=mRNA /PROD=Similar to eukaryotic translation initiationfactor 3, subunit 8 (110kD) //DB XREE=qi:12653522 /UG=HS.4835 eukaryotic translation initiation factor 3. subunit β (110kD) /FFi=ch·scionce3 1 |
| 213142_x_at | 0:AV700415 /FEA-EST /DB_XREF=gi:10302386 /DB_XREF=est:AV700415 /CLONE=GKCDGA04 /UG=Hs.12969 hyp |
| 217817_at | ricludes gb: BE891920 /FEA=EST /DB XREF=gi:10351728 /DB XREF=est:601435490F |
| 216041_x_at | Process 2 Compres, sur 148.1 / DEF=Homo sapiens |
| | /FEA=mRNA /DB XREF=gi:10435243 /UG=Hs.180577 granulin |
| 20268/_s_at | gb:U>/U>9.1 /DEF=Homo sapiens Apo-2 11gand mRNA, complete cds. /FEA=mRNA /PROD=Apo-2 ligand /DB XREF=gi:1336207 /UG=Hs.83429 tumor necrosis factor (ligand) superfamily, member 10 /FL=qb:U37518.1 db:U57059.1 db:NM 003810 1 |
| 215230_x_at | 227 /DB_XREF=est:ah15e04.sl /CLONE=IMAGE:1156 subunit 8 (110kp) |
| 200647_x_at | initiation factor 3, subunit 8 (1 |
| | ion initiation factor 3, subunit 8 (110kD) /FL=gb: |
| 200078_s_at | gb:BC005876.1 /DEF=Homo sapiens, ATPase, H+ transporting, lysosomal (vacuolar proton pump) 21kD, clone MGC:4498, mRNA, complete cds. /FEA=mRNA /PROD=ATPase, H+ transporting, lysosomal (vacuolarproton pump) 21kD /DB_XREF=gi:13543437 //FL=qb:BC005876.1 |
| 202169_s_at | gb:RF302110.1 /DEF=Homo sapiens alpha-aminoadipic semialdehyde dehydrogenase-phosphopantetheinyl transferase mRNA, complete cds. /FEA=mRNA /PROD=alpha-aminoadipic semialdehydedehydrogenase-phosphopantetheinyl transferase |
| | -Hs.64595 aminoadipate-semialdehyde dehydrogenase-phosphopantetheinyl t 86978.1 gb:AF151838.1 gb:AF151057.1 gb:NM 015423.1 gb:AF201943.1 |
| 202659_at | 801.1 /DEF-Homo sapiens proteasome (prosome, macropain) subunit, 10 /PROD-proteasome (prosome, macropain) subunit, betatype, 10 /DE |
| ١ | * |
| 271488_S_at | gp:ArzJoyz4.1 / DEr=Homo sapiens brain acetylcholinesterase putative membrane anchor mRNA, complete cds. /FEA=mRNA /PROD=brain acetylcholinesterase putative membraneanchor /DB_XREF=gi:7341254 /UG=Hs.107187 divalent cation tolerant protein CUTA /FL=gb:AF230924.1 |
| 221622_s_at | gb:AF246240.1 /DEF=Homo sapiens HT026 mRNA, complete cds. /FEA=mRNA /PROD=HT026 /DB_XREF=gi:12005514 //UG=Hs:24371 uncharacterized hypothalamus protein HT007 /FL=gb:AF246240.1 |

| 206050_s_at | sapiens ribonucleaseangiogenin inhibitor (RNH), mRNA. /FEA=mRNA /GEN=RNH nin 06564 /UG-Hs.75108 ribonucleaseangiogenin inhibitor /FL=gb:M36717.1 gb:NM 002939. |
|-------------|---|
| 212199_at | 66962 /FEA=EST /DB_XREF=gi:12919867 /DB_XREF=est:AL566962 /CL ive ribosomal protein S1 mRNA |
| 201861_s_at | 65566 /FEA-EST /DB_XREF-gi:12332781 /DB_XREF-est:602277032F1 /C h repeat (in FLII) interacting protein 1 /FI-gb:NM 004735.1 |
| 212795_at | Consensus includes gb:AL137753.1 /DEF-Homo sapiens mRNA; cDNA DKFZp434K1412 (from clone DKFZp434K1412). /FEA=mRNA /DB XREF=gi:6808455 /UG-Hs.12144 K1AA1033 protein |
| 213574_s_at | Consensus includes gb:AA861608 /FEA=EST /DB_XREF=gi:2953748 /DB_XREF=est:ak34e01.s1 /CLONE=IMAGE:1407864 /UG=Hs.180446 karyopherin (importin) beta 1 |
| 215171_s_at | Consensus includes gb:AK023063.1 /DEF-Homo sapiens cDNA FLJ13001 fis, clone NT2RP3000341, highly similar to Homo sapiens mitochondrial inner membrane preprotein translocase Timl7a mRNA, nuclear gene encoding mitochondrial protein. /FEA-mRNA /DB XREF-gi:10434808 /UG-Hs.20716 translocase of inner mitochondrial membrane 17 (yeast) homolog A |
| 210137_s_at | |
| 212943_at | /DEF=Homo sapiens mRNA for KIAA0528 protein, partial cds. /FEA=mRNA /GEN=KIAA05 gi:6683714 /UG=Hs.30656 KIAA0528 gene product |
| at | CALIFP (CALIF) mRNA, complete cds. /FEA=mRNA /GEN=CALIF /PROD=CALIFP / |
| 209005_at | 'DEF-Homo sapiens p45SKP2-like protein mRNA, complete cds. /FEA-mRNA /PRO 18696 /UG-Hs.5548 f-box and leucine-rich repeat protein 5 /FI-gb:AF199420 |
| 202899_s_at | gb:NM 003017.1 /DEF=Homo sapiens splicing factor; arginineserine-rich 3 (SFRS3), mRNA. /FEA=mRNA /GEN=SFRS3 /PROD=splicing factor, arginineserine-rich 3 /DB_XREF=gi:4506900 /UG=Hs.167460 splicing factor, arginineserine-rich 3 /FT=Ab:1.10838 1 ab:NM 003017 1 |
| 208777 s at | apiens 26S proteasome subunit 9 mRNA, co |
| | 50045 /UG=Hs.90744 7.1 gb:BC004430.1 c |
| 219889_at | |
| | /UG-Hs.126057 frequently rearranged in advanced T-cell lymphomas /FL=gb:U58975.2 gb:NM 005479.1 |
| 204961_s_at | sapiens neutropint cycosoffe factor 1 /DR YBFFEGI-45K1784 /NG=Ne 1583 neutrophil cytosolic facto |
| | cous disease, autosomal 1) /FIL-qb:BC002816.1 gb:AF330627.1 gb:M55067.1 gb:N |
| 217106_x_at |)91078.1 /DEF-Homo sapiens clone 559 unknown mRNA, complete sequence. /FEA=mRNA .:3859993 /UG-Hs.125819 putative dimethyladenosine transferase |
| 212224_at | irogenase 1, soluble (ALD .76392 aldehyde dehydrog |
| 204125_at | |
| 217752_s_at | sapiens hypothetical protein FLJ10830 (FLJ10830), mRNA. /FEA-mRNA /GEN-FLJ10830 in 2698 /UG-Hs.273230 hypothetical protein FLJ10830 /FL-gb:BC001375.1 gb:BC003176.1 gb:NN |
| 221087_s_at | gb:NM_014349.1 /DEF=Homo sapiens apolipoprotein L, 3 (APOL3), mRNA. /FEA=mRNA /GEN=APOL3 /PROD=apolipoprotein L, 3 |

| | /DB_XREF=gi:7656972 /UG=Hs.241535 apolipoprotein L, 3 /FL=gb:AY014905.1 gb:AF070675.1 gb:NM_014349.1 |
|-------------|---|
| 218139_s_at | gb:NM 018229.1 /DEF=Homo sapiens hypothetical protein FLJ10813 (FLJ10813), mRNA. /FEA=mRNA /GEN=FLJ10813 /PROD=hypothetical protein FLJ10813 /DB_XREF=gi:8922687 /UG=Hs.106210 hypothetical protein FLJ10813 /FL=gb:AL136685.1 gb:NM 018229.1 |
| 213878_at | Consensus includes gb:AI685944 /FEA=EST /DB_XREF=g1:4897238 /DB_XREF=est:tu38g02.x1 /CLONE=INAGE:2253362 /UG=Hs.235069 RecQ protein-like (DNA helicase Q1-like) |
| 201346_at | gb:NM 024551.1 /DEF-Homo sapiens hypothetical protein FLJ21432 (FLJ21432), mRNA. /FEA-mRNA /GEN-FLJ21432 /PROD-hypothetical protein FLJ21432 /DB_XREF-gi:13375714 /UG-Hs.11641 hypothetical protein FLJ21432 /FL-gb:NM 024551.1 gb:BC004906.1 |
| 206874_s_at | 38761 /DEF=Human DNA sequence from clone RP11-16H23 |
| × - | Contains the gene KIAA0204 (HSLK) for a protein kinase, the COL17A1 gene for collagen type XVII alpha 1 (BP180), ESTs and GSSs /FEA=mRNA 2 /DB_XREF=gi:8573811 /UG=Hs.105751 Ste20-related serinethreonine kinase |
| 221808 at | Consensus includes ab:NM 004251.1 / DEF=Homo sabiens RAB9. member RAS oncodene family (pand) menn / TEXCDS /CENnap. |
| l | loogene family /DB_XREF=gi:4759011 /UG=Hs.28726 RAB9, member RAS oncogene family /1251.1 |
| 211138_s_at | /DEF-Homo sapiens, Similar to kynurenine 3-monooxygenase (kynurenir |
| | complete cds. /FEA=mRNA /PROD=Similar to kynurenine 3-monooxygenase(kynurenine 3-hydroxylase) /DB_XREF=gi:13529016 /UG=Hs.107318 kynurenine 3-monooxygenase (kynurenine 3-hydroxylase) /FL=ob:RC005297 1 |
| 201386 s at | sapiens dead box protein 15 mRNA, complete |
| 1. | Hs.5683 DEADH (Asp-Glu-Ala-AspHis) box polypeptide 15 /FL-gb:AB001636.1 gb:NM 001 |
| 203037_s_at | gb:NM_014751.1 /DEF=Homo sapiens KIAA0429 gene product (KIAA0429), mRNA. /FEA=mRNA /GEN=KIAA0429 /PROD=KIAA0429 gene |
| | product /DB XREE=gi:7662113 /UG=Hs.77694 KIAA0429 gene product /FL=qb:AB007889.1 qb:NM 014751.1 |
| 218356_at | cell division protein FtsJ (FJH1), mRNA. |
| 204405_x_at | sapiens |
| | /PROD=putative dimethyladenosine transferase /DB_XREF=g1:7657197 /UG=Hs.125819 putative dimethyladenosine transferase /FL=gb:AF102147.1 gb:NM 014473.1 |
| 201608_s_at | ear phosphoprotein similar to S. cerevisiae PWP1 (PWP1), mRt to S. cerevisiaePWP1 /DB XREF=gi:5902033 |
| 201569 s at | gb:NM 015380.1 /DEF=Homo sapiens CGI-51 protein (CGI-51), mRNA, /FEA=mRNA /GRN=CGI-51 /PROD=CGI-51 protein |
| | 80.1 |
| 222244_s_at | Consensus includes gb:AK000749.1 /DEF=Homo sapiens cDNA FLJ20742 fis, clone HEP06891. /FEA=mRNA /DB_XREF=gi:7021031 //GG=Hs.52184 hypothetical protein FLJ20618 |
| 201241_at | gb:NM 004939.1 /DEF-Homo sapiens DEADH (Asp-Glu-Ala-AspHis) box polypeptide 1 (DDX1), mRNA, /FEA-mRNA /GEN-DDX1 //PROD-DEADH (Asp-Glu-Ala-AspHis) box polypeptide 1 /DB XREF-q1:4826685 /UG-Hs.78580 DEADH (Asp-Glu-Ala-AspHis) hox |
| | |
| 220731_s_at | gb:NM 018090.1 /DEF-Homo sapiens hypothetical protein FLJ10420 (FLJ10420), mRNA. /FEA-mRNA /GEN-FLJ10420 /PERP // PROD-hypothetical protein FLJ10420 /DB XREF-qi:8922415 /UG-Hs.289087 hypothetical protein FLJ10420 /Fim-h.NM 018090 1 |
| 208799_at | oteasome (prosome, macropain) subunit, beta type, 5, clone MGC:2175, mRNA, |

| | /FEA=mRNA /PROD=proteasome (prosome, macropain) subunit, betatype, 5 /DB XREF=gi:13278740 /UG=Hs.78596 proteasome (prosome, macropain) subunit, beta type, 5 /FL=gb:BC004146.1 gb:NM 002797.1 gb:D29011.1 |
|-------------|---|
| 221718_s_at | pe II CAMP-dependent protein kinase (Ht31) mRNA, complete o XREF=gi:184434 /FL-gb:M90360.1 |
| 208662_s_at | /FEA=EST /DB_XREF=g1:5590502 /DB_XREF=est:w192e09.x1 e_repeat_domain 3 /FL=gb:D84294.1 |
| 202118_s_at | 541758 /FEA=EST /DB_XREF=qi:2288192 /DB_XREF=est:ni87d05.s1 /C: /FL=qb:AB014536.1 qb:NM 003909.1 qb:AF077226.2 |
| 201740_at | gb:NM 004551.1 /DEF=Homo sapiens NADH dehydrogenase (ubiquinone) Fe-S protein 3 (30kD) (NADH-coenzyme Q reductase) (NDUFS3), mRNA. /FEA-mRNA /GEN-NDUFS3 /PROD-NADH dehydrogenase (ubiquinone) Fe-S protein 3(30kD) (NADH-coenzyme Q reduc- |
| | tase) /DB_XREF=gi:4758787 /UG=Hs.5273 NADH dehydrogenase (ubiquinone) Fe-S protein 3 (30kD) (NADH-coenzyme Q reductase) /FL=gb:BC000617.1 gb:AF067139.1 gb:NM 004551.1 gb:AF100743.1 |
| 215952_s_at | |
| 212796_s_at | 95608 /FEA=EST /DB_XREF=gi:11082676 /DB_XREF=est:7n85f03.x1 /CLONE=IMAGE:357 otein |
| 210561_s_at | gb:ALI10243.1 /DEE=Homo sapiens mRNA; cpNA DKEZp56480482 (from clone DKEZp56480482); complete cds. /FEA=mRNA |
| | |
| 202220_at | sapiens KIAA0907 protein (KIAA0907), mRNA. /FEA=mRNA /GEN=KIAA0907 /PROD=KIAA0907 -Hs.24656 KIAA0907 protein /FL=gb:AB020714.1.gb:NM 014949.1 |
| 203739_at | gb:NM_006526.1 /DEF=Homo sapiens zinc finger protein 217 (ZNF217), mRNA. /FEA=mRNA /GEN=ZNF217 /PROD=zinc finger protein |
| | XREF=gi:5730123 /UG=Hs.155040 zinc finger protein 217 /FL=gb:AF041259.1 gb:NM 006526.1 |
| 211036_x_at | apiens, anaphase-promoting complex subunit 5, clone MGC:13295, mRNA, complete cds complex subunit 5 /DB XREF=gi:13623410 /FL=gb:BC006301.1 |
| 204060_s_at | gb: NN 005044.1 /DEF=Homo sapiens protein kinase, X-linked (PRKX), mRNA. /FEA=mRNA /GEN=PRKX /PROD=protein kinase, X- linked |
| | Is.147996 protein kinase, X-linked /FL-gb:NM 005044.1 |
| 64064_at | Cluster Incl. A1435089:th95b11.x1 Homo sapiens CDNA, 3 end /clone=IMAGE-2126397 /clone_end=3 /gb=A1435089 /gi=4300737 //ug=Hs.26194 /len=891 |
| 201515_s_at | sapiens translin (TSN), mRNA. /EEA=mRNA /GEN=TSN /PROD=translin /D .=qb:NM 004622.1 |
| 219093_at | sapiens hypothetical protein FLJ20701 (FLJ20701), mRNA. /FEA-mRNA /GEN-FLJ2070 n FLJ20701 /DB XREF-gi:8923631 /UG-HS.169764 hypothetical protein FLJ20701 /FL |
| 221739_at | 24093 /FEA=EST /DB_XREF=gi:12787586 /DB_XREF=est:AL524093 /CLONE=CS0 protein EUROIMAGE1875335 |
| 208822_s_at | /FEA=mRNA /PROD=i :ein 3 /FL=gb:U18 |
| 218160_at | gb:NM 014222.1 /DEF=Homo sapiens NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 8 (19kD, PGIV) (NDUFA8), mRNA. /FEA=mRNA /GEN=NDUFA8 /PROD=NADH dehydrogenase (ubiquinone) 1 alphasubcomplex, 8 (19kD, PGIV) /DB XREF=gi:7657368 |
| • | genase (ubiquinone) l alpha subcomp |
| 221728_x_at | Consensus includes gb:AA628440 /FEA=EST /DB_XREF=gi:2540827 /DB_XREF=est:af26f02.s1 /CLONE=IMAGE:1032795 /UG=Hs.83623 nuclear receptor subfamily 1, group I, member 3 |

| 210943_s_at | gb:U84744.1 /DEF=Human Chediak-Higashi syndrome protein short isoform (LYST) mRNA, complete cds. /FEA=mRNA /GEN=LYST /PROD=Chediak-Higashi syndrome protein short isoform /DB XREF=gi:2654473 /MG=Hs 36508 Chediak-Higashi syndrome |
|-------------|--|
| ŀ | THE STATE OF THE S |
| 209104_s_at | gb:BC000009.1 /DEF=Homo sapiens, likely homolog of yeast Nhp2, component of the HACA snoRNP; hypothetical protein |
| | C:1038, mRNA, complete cds. /FEA-mRNA /PROD=likely homolog of yeast Nhp2, component |
| | nypounetical procein fiologgy / UB_AREE=g1:12052540 / UG≃HS.23990 nucleolar protein family A, member 2 (HACA small nucleo lar RNPS) |
| - | /FL=gb:BC00009.1 gb:NM 017838.1 |
| 217987_at | gb:NM_019048.1 /DEF=Homo sapiens hypothetical protein (FLJ20752), mRNA. /FEA=mRNA /GEN=FLJ20752 /PROD=hypothetical pro- |
| | DB XREF=gi:9506696 /UG=Hs.101364 hypothetical protein /FL=qb:BC001243.1 qb:NM 019048.1 |
| 201699_at | 06.1 /DEF-Homo sapiens proteasome (prosome, macropain) 26S subu |
| : | |
| | subunit, ATPase, |
| 201359_at | /DEF=Homo sapiens coatomer protein complex, subunit beta (COPB), mRNA. /F |
| : | /PROD=coatomer protein complex, subunit beta /DB_XREF=qi:7705368 /UG=Hs.3059 coatomer protein complex, subunit beta. /FL=qb:AF084457.1 qb:AL136593.1 qb:NM 016451.1 |
| 210027_s_at | urinic endonuclease (APE) mRNA, complete cds. /FEA=mRNA / |
| 202096 s at | /UG-ns.13/22 Arra Huttease (Multilunctional DNA repair enzyme) /FL=gb:BCU04979.1 gb:N80261.1 |
| : | |
| | transcript variant PBR, mRNA. /FEA=mRNA /GEN=BZRP /PROD=peripheral benzodiazapine receptor /DB_XREF=gi:6382068 /UG=Hs.202 benzodiazapine receptor (peripheral) /FI=qb:BC001110 1 db:M36035 1 db:NM 000714 2 |
| 214948_s_at | 50136.1 /DEF=Homo sapiens mRNA; cDNA DKFZp586L141 (from clone DKFZp5 |
| 200042 at | 105 AREE-91:4001310 (00-13:14031) REPORT REPORT TO THE DRA DRESPONDENTAL (IFOM CIONE DRESDSWELLAL) |
| | 55.5 E. D. T. C. |
| ٠, | JD:AF161466.1 gD:NM 014306.1 gb:AF155658.1 |
| בוספסק ב | ٠. |
| 200041_s_at | M_004640.1 /DEF=Homo sapiens HLA-B associated transcript-1 (D6S81E), mRNA. /FEA=mRNA /GEN=D6S81E / |
| | dreu transcript-1 /DB XREF=qi:4758111 /UG=Hs.55296 HLA-B associated transcript-1 /FL=qb:Rc004350 1 qb:NM 004640 1 |
| 216321_s_at | 348.1 /DEF=Human mRNA for beta-glucocorticoid receptor (clone OB10). /FEA=mRNA |
| 214574 x at | 72 nuclear receptor subfamily 3, group |
| :¦ | or, poly (D6849E), mRNA. /FEA=CDS /GEN=D6849E |
| 205370 x at | HS.00411. Lymphocyte antigen 11/ /Filego:NM UU/161.1 |
| | ple syrup urine disease) (DBT), mRNA. /FEA=mRNA /GEN=DBT |
| | ecomplex |
| | omplex; maple syrup urine |
| 213872_at | /DB_XREF=est:hv76g09.x1 /CLONE= |

| | - |
|-------------|--|
| | |
| 201576_s_at | sapiens galactosidase, beta 1 (GLB1), mRNA. / 1965 /UG=Hs.79222 galactosidase, beta 1 /FI=gb |
| 202548_s_at | sapiens PAK-interacting exchange factor beta (PBSSFR), mkn4. /FEA=mknA /GEN=PBSSSFR Change factor beta /DB_XREF=gi:4505572 /UG=Hs.172813 PAK-interacting exchange facto 1899.1 |
| 204028_s_at | =Homo sapien CENA /PROD=r TPase activa |
| 202435_s_at | 154504 /FEA=EST /DB_XREF=gi:11016025 /DB_XREF=est:AU15450 P450, subfamily I (dioxin-inducible), polypeptide 1 (gla 3688.1 |
| 217980_s_at | <pre>sapiens hypothetical protein FLJ20484 (FLJ20484), mRNA. /FEA=mRNA /GEN= in FLJ20484 /DB XREF=q1:8923447 /UG=Hs.5080 hypothetical protein FLJ2048)1040.1 gb:NM 017840.1 gb:AF183428.1</pre> |
| 201112_s_at | sapiens chromosome segregation 1 (yeast homolo bme segregation 1 (yeast homolog)-like /DB_XREE -gb:U33286.1 gb:AF053641.1 gb:NM 001316.1 |
| 202322_s_at | sapiens geranylgeranyl diphosphate synthase l (GGPSI), mRNA. /FEA-mRNA /GEN=GGPS1 nosphate synthase l /DB XREF=gi:4758429 /UG=Hs.55498 geranylgeranyl diphosphate synthase l 05252.1 gb:AB017971.1 gb:AB016043.1 gb:AB019036.1 gb:NN 004837.1 gb:AF125394.1 |
| 203545_at | s hypothetical protein MGC2840 similar to a putative glucosyltransferase pothetical protein MGC2840 similar to aputative glucosyltransferase /DB-ein MGC2840 similar to a putative glucosyltransferase /FI=gb:BC001133.1 |
| 212646_at | gb:D42043.1 1084 protein |
| 212591_at | gb:AA887480 / ike protein |
| 218005_at | |
| 201165_s_at | <pre>gb.BE670915 /FEA=EST /DB XREF=gi:10031456 /DB XREF=est:7e43a09.x1 io (Drosophila) homolog I /FL=gb:AF315592.1 gb:NM 014676.1</pre> |
| 212239_at | /FERA=EST / DB XREF=gi:4990374 / DB XREF=est:tw66a10.X1 / C. kinase, regulatory subunit, polypeptide 1 (p85 alpha) |
| 204386_s_at | |
| 214853_s_at | /FEA=EST /DB XREF=gi:3430138 /DB XREF= 2 domain-containing) transforming prot |
| 210580_x_at | uman estrogen sulfotransferase mRNA, complete cds. /FEA=mRNA /PROD=estrogen sulfotra //UG=Hs.274614 sulfotransferase family, cytosolic, 1A, phenol-preferring, member 3 / |
| 222150_s_at | lone LNG0/3/9, highly similar to HSTU: -gi:10439670 /UG-Hs.12969 hypothetical |
| 209265_s_at | gb:BC001650.1 /DEF=Homo sapiens, Similar to putative methyltransferase, clone MGC:2708, mRNA, complete cds. /FEA=mRNA /PROD=Similar to putative methyltransferase /DB_XREF=gi:12804482 /UG=Hs.268149 putative methyltransferase /FL=gb:BC003031.1 gb:BC001650.1 gb:NM 019852.1 |

| 218807_at | gb:NM 006113.2 /DEF=Homo sapiens vav 3 oncogene (VAV3), mRNA. /FEA=mRNA /GEN=VAV3 /PROD=vav 3 oncogene /DB XREF=gi:7262390 |
|-------------|--|
| | gene /FL=gb:AF067817.1 gb:AF118887.1 gb:NM 006 |
| 209043_at | gb:RF033026.1 /DEF=Homo sapiens bifunctional ATP sulfurylaseadenosine 5-phosphosulfate kinase mRNA, complete cds. /FEA=mRNA /PROD=bifunctional ATP sulfurylaseadenosine5-phosphosulfate kinase /DB_XREF=gi:3378100 /UG=Hs.3833 3-phosphoadenosine 5-phosphosulfate synthase 1 /FL=cb:3F033076 1 ch:3F016496 1 ch:NW 005443 1 ch:aF105222 1 |
| 219777_at | .1 / DEF=Homo sapiens hypothetical protein FLJ22690 (FLJ22690), mRNA. /FEA=mRNA /GEN=FLJ22690 etical protein FLJ22690 /DB XREF=qi:13376008 /UG=Hs.105468 hypothetical protein FLJ22690 /FL=c |
| 218048_at | |
| 201986_at | Consensus includes gb: AB011165.1 /DEF=Homo sapiens mRNA for KIAA0593 protein, partial cds. /FEA=mRNA /GEN=KIAA0593 /PROD=KIAA0593 protein /DB XREF=gi:3043709 /UG=Hs.11861 thyroid hormone receptor-associated protein, 240 kDa subunit |
| 219799_s_at | |
| 208688_x_at | gp:078525.1 /DEF-Homo sapiens eukaryotic translation initiation factor (eIF3) mRNA, complete cds. /FEA-mRNA /GEN-eIF3 /PROD-eukaryotic translation initiation factor /DB_XREF-gi:2558667 /UG-Hs.57783 eukaryotic translation initiation factor |
| | 3, subunit 9 (eta, 116kD) /FI=gb:U78525.1 |
| 200001_at | gb:NM_001749.1 /DEF=Homo sapiens calpain 4, small subunit (30K) (CAPN4), mRNA. /FEA=mRNA /GEN=CAPN4 /PROD=calpain 4 small subunit |
| 218118_s_at | translocate of inner mitochondri nner mitochondrial membrane 23 (ye r mitochondrial membrane 23 (yea |
| 201840_at | apiens neural precursor cell expressed, demonstrated expressed, demonstrated expressed, developmentally |
| 38710_at | mo sapiens mRNA; cDNP /len=1904 |
| 212287_at | Consensus includes gb:BF382924 /FEA=EST /DB_XREF=gi:11364313 /DB_XREF=est:601816985F1 /CLONE=IMAGE:4050909 /UG=Hs.197803 KIAA0160 protein |
| 203098_at | .1 /DEF=Homo sapiens /PROD=hypothetical p |
| 218905 at | AF081258.1 gb:AF081259.1 gb:NM 004824.1 sabiens hypothetical protein FLJ20530 (FLJ20530). mRNA. /FER=mRNA /GEN=FLJ2053 |
| 218303 x at | cal protein FLJ20530 / DB XREE-g1:8923495 /UG-HS.279521 hypothetical protein /DEF=Homo sapiens hypothetical protein (LOC51315), mRNA. /FEA=mRNA /GEN=LOC5 |
| 208705_s.at | tein /DB XREF-gi:7706155 /UG-Hs.5721 hypothetical protein /FL-gb:AF208845.1 gb:AF217520.1 gb:NM 016618.1 Consensus includes gb:BG481972 /FEA=EST /DB XREF=gi:13414251 /OB XREF=est:602526894F1 /CLONE=IMAGE:4650406 /UG-Hs.286236 enkarvotic translation initiation factor 5 /FT-ch:AT080107 1 |
| 218715_at | sapiens hepatocellular carcinoma cinoma-associated antigen 66 /DB inco 66 /FE=ch: AF744135 1 ch: AF11 |
| 202184_s_at | sapiens hypothetical protein FLJ10814 (|

| | /PROD=hypothetical protein FLJ10814 /DB_XREF=gi:8922689 /UG=Hs.12457 hypothetical protein FLJ10814 /FL=gb:NM_018230.1 |
|--------------------|---|
| 202004_x_at | <pre>gb:NM_003001.2 /DEF=Homo sapiens succinate dehydrogenase complex, subunit C, integral membrane protein, 15kD (SDHC), nuclear gene encoding mitochondrial protein, mRNA. /FEA=mRNA /GEN=SDHC /PROD=succinate dehydrogenase complex, subunit Cprecursor /DB_XREF=g1:9257243 /UG=Hs.3577 succinate dehydrogenase complex, subunit C, integral membrane protein, 15kD /FL=gb:U57877.1 gb:NM 003001.2</pre> |
| 209435_s_at | gb:BC000265.1 /DEF=Homo sapiens, clone MGC:3182, mRNA, complete cds. /FEA=mRNA /PROD=Unknown (protein for MGC:3182) /DB XREF=gi:12653008 /UG=Hs.315417 Homo sapiens, clone MGC:3182, mRNA, complete cds /FL=gb:BC000265.1 |
| 221044_s_at | gb:NM 021616.1 /DEF=Homo sapiens ring finger protein 21, interferon-responsive (RNF21), mRNA. /FEA-mRNA /GEN-RNF21 /PROD=ring finger protein 21, interferon-responsive /DB_XREF=gi:11055977 /UG=Hs.125300 ring finger protein 21, interferon-responsive /FL=gb:AB039903.1 gb:NM 021616.1 |
| 200992_at | Consensus includes gb:AL137335.1 /DEF=Homo sapiens mRNA; cDNA DKF2p434A179 (from clone DKF2p434A179); partial cds. /FEA=mRNA /GEN=DKF2p434A179 /PROD=hypothetical protein /DB_XREF=gi:6807827 /UG=Hs.5151 RAN binding protein 7 /FL=gb:AF098799.1 gb:NN 006391.1 |
| 201900_s_at | gb:NM 006066.1 /DEF=Homo sapiens aldo-keto reductase family 1, member Al (aldehyde reductase) (AKRIA1), mRNA. /FEA=mRNA /CEN=AKRIA1 /PROD=aldo-keto reductase family 1, member Al (aldehyde reductase) /DB XREF=gi:5174390 //CEA-RS.89529 aldo-keto reductase family 1, member Al (aldehyde reductase) /FL=gb:BC000670.1 gb:J04794.1 gb:NM 006066.1 |
| 202560_s_at | sapiens DKFZP547E1010 protein (DKFZP547E1010), m in /DB XREF-gi:7661589 /UG-Hs.323817 DKFZP547E10 |
| 201105_at | sapiens lectin, galactoside-binding, soluble, 1 (galectin 1) (LGALS1), m alactosidase binding lectin precursor /DB_XREF=gi:6006015 /UG-Hs.227751 1 ectin 1) /FL=gb:BC001693.1 gb:J04456.1 gb:NM 002305.2 |
| 204546_at | gb: NW 014732.1 /DEF=Homo sapiens KIAA0513 gene product (KIAA0513), mRNA, /FEA=mRNA /GEN=KIAA0513 /PROD=KIAA0513 gene product /DB XREF=gi:7662163 /UG=Hs.301658 KIAA0513 gene product /FL=gb:AB011085.1 gb:NN 014732.1 |
| 214181_x_at | Consensus includes gb:AI735692 /FEA=EST /DB_XREF=gi:5057216 /DB_XREF=est:at20b12.x1 /CLONE=IMAGE:2355647 /UG=Hs:88411 lymphocyte antigen 117 |
| 219032_x_at | gb:NM 014322.1 /DEF-Homo sapiens opsin 3 (encephalopsin) (OPN3), mRNA. /FEA-mRNA /GEN-OPN3 /PROD-opsin 3 (encephalopsin) /DB XREF-gi:7657070 /UG-Hs.279926 opsin 3 (encephalopsin) /FL-gb:AF140242.1 gb:NM 014322.1 |
| 205306_x_at | 3-4145 /FEA-EST /DB_XREF=gi: 3-monooxygenase (kynurenine |
| 202837_at | db:NM 006700.1./DEF=Homo sapiens FIN29 gene product (FIN29), mRNA. /FEA=mRNA /GEN=FIN29 /PROD=FIN29 gene product /DB XREF=gi:5729827 /UG=Hs.5148 FIN29 gene product /FL=gb:BC003553.1 gb:AB007447.1 gb:NM 006700.1 |
| 202153_s_at | sapiens hypothetical protein (DKF2p5 n /DB XREF=gi:7705354 /UG=Hs.9877 hy |
| 209341_s_at | |
| 212493_s_at | 761110 /F |
| 201600 at | <pre>gb:NM 007273.1 /DEF=Homo sapiens B-cell associated protein (REA), mRNA. /FEA=mRNA /GEN=REA /PROD=B-cell associated pro- tein /DB_XREF=gi:6005853 /UG=Hs.7771 B-cell associated protein /FL=gb:AF150962.1 gb:NM_007273.1 gb:AF126021.1 gb:AF178980.1</pre> |
| 201144 <u>s</u> at | <pre>gb:NM_004094.1 /DEF=Homo sapiens eukaryotic translation initiation factor 2, subunit 1 (alpha, 35kD) (EIF2S1), mRNA. /FEA=mRNA /GEN=EIF2S1 /PROD=eukaryotic translation initiation factor 2, subunit 1 (alpha, 35kD) /DE_XREF=gi:4758255 /UG=Hs.151777 eukaryotic translation initiation factor 2, subunit 1 (alpha, 35kD) /FL=gb:BC002513.1 gb:J02645.1 gb:NM 004094.1</pre> |
| 211985_s_at | Consensus includes gb:AI653730 /FEA=EST /DB_XREF=gi:4737709 /DB_XREF=est:wb36f12.xl /CLONE=IMAGE:2307791 /UG=Hs.279009 matrix Gla protein |

| 00000 | |
|-------------|---|
| 701220 at | <pre>gb:nd_U05U04.1 /DEF=Homo sapiens NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 8 (19kD, ASHI) (NDUFB8), mRNA. //FEA=mRNA /GEN=NDUFB8 /PROD=NADH dehydrogenase (ubiquinone) 1 betasubcomplex, 8 (19kD, ASHI) /DB_XREF=gi:4826853 //UG=Hs.198273 NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 8 (19kD, ASHI) //FI=gb:BC000466.1 gb:AF044958.1 gb:AF077028.1 gb:NM 005004.1 gb:AL080056.1</pre> |
| 208828_at | gb:BC004170.1 /DEF=Homo sapiens, histone fold protein CHRAC17; DNA polymerase epsilon p17 subunit, clone MGC:2725, mRNA, complete cds. /FEA=mRNA /PROD=histone fold protein CHRAC17; DNA polymeraseepsilon p17 subunit /DB_XREF=gi:13278800 /UG=HS.108112 histone fold protein CHRAC17; DNA polymerase epsilon p17 subunit /FL=gb:BC003166.1 gb:BC004170.1 gb:AF226077.1 gb:NM 017443.1 |
| 204020_at | Consensus includes gb:BF739943 /FEA=EST /DB_XREF=g1:12066607 /DB_XREF=est:7041b04.x1 /CLONE=IMAGE:3576534 /UG=Hs.29117 purine-rich element binding protein A /FL=qb:M96684.1 db:NM 005859.1 |
| 217942_at | sapiens MDS023 protein (MDS023), mRNA. /FF -Hs.10724 MDS023 protein /Fri=gb:AF182422.1 |
| 203514_at | Consensus includes gb:BF971923 /FEA=EST /DB_XREF=qi:12339138 /DB_XREF=est:602240326F1 /CLONE=IMAGE:4328791 /UG=Hs.29282 mitogen-activated protein kinase kinase 3 /FE=qb:U78876.1 db:NM 002401.1 |
| 204334_at | /DB_XREF=g1:2216103 /DB_XREF=est:aD40d10.s1 quitous) /FL=gb:AB01513Z.1 qb:NM 003709.1 |
| 203531_at | 135809 /FEA=EST /DB_XREF=91:11448124 /DB_XF :L=gb:AF327710.1 gb:AF017061.1 gb:NM 003476 |
| 205022_s_at | gb:NM_005197.1 /DEF=Homo sapiens checkpoint suppressor 1 (CHES1), mRNA. /FEA=mRNA /GEN=CHES1 /PROD=checkpoint suppressor 1 /DE XREF=g1:4885136 /UG=Hs.211773 checkpoint suppressor 1 /FL=qb:U68723.1 qb:NM 005197.1 |
| 201532_at | /pe, 3 (PSMA3), mRNA. / |
| 100000 | nnit, alpha type, 3 /FL=gb:BC005265.1 gb:NM 002788.1 |
| | |
| 4- 000000 | 817.1 |
| 203200_at | Consensus includes gb:AL536517 /FEA-EST /DB_XREF=gi:12800010 /DB_XREF=est:AL536517 /CLONE=CS0DF038YH13 (5 prime) /UG=Hs.78995 MADS box transcription |
| 11- | eptide C (myocyte enhancer factor 2C) /FL=gb:L06 |
| 21/945_at | gb:NM_025238.1 /DEF=Homo sapiens BTB (POZ) domain containing 1 (BTBD1), mRNA, /EEA=mRNA /GEN-BTBD1 /PROD=BTB (POZ) domain containing 1 /DB XREF=qi:13376847 /UG=HS.21332 BTB (POZ) domain containing 1 /FI=ch-1136863 1 |
| | 238.1 gb:AF355402.1 |
| 201491_at | FEA=mRNA /GEN=C140RF3 |
| | .1 gb:NM 012111.1 gb:AF164791 |
| 204868_at | gb:NM 001545.1 /DEF=Homo sapiens immature colon carcinoma transcript 1 (ICT1), mRNA. /FEA=mRNA /GEN=ICT1 //DB XRFF==i-4 //OFTAFA |
| | 5.1 |
| 217826_s_at | gb:NM 016021.1 /DEF=Homo sapiens CGI-76 protein (LOC51632), mRNA. /FEA=mRNA /GEN=LOC51632 /PROD=CGI-76 protein /DB XREF=gi:7706311 /UG=Hs.184325 CGI-76 protein /FI=qb:AF151834.1 db:AF161502.1 db:AF151039.1 db:MM 01603.1 |
| 212534_at | 56 /CLONE=HEMBA1000798 |
| 211284_s_at | apiens, Similar 3114 /UG=Hs.1809 |
| 212188_at | 51075 /FEA=EST /DB_XREF=gi:23 |
| | |

| 201458_s_at | gb:NM 004725.1 /DEF=Homo sapiens BUB3 (budding uninhibited by benzimidazoles 3, yeast) homolog (BUB3), mRNA. /FER=mRNA //GEN=BUB3 /PROD=BUB3 (budding uninhibited by benzimidazoles 3, yeast) homolog /DB XREF=gi:4757879 /UG=Hs.40323 BUB3 //DD //DD APASIMIPITED by benzimidazoles 3, yeast) homolog /FI=Hb\RCOORT38 1 gb\rackstructure by benzimidazoles 3, yeast) homolog /FI=Hb\RCOORT38 1 gb\rackstructure by benzimidazoles 3, yeast) homolog /FI=Hb\RCOORT38 1 gb\rackstructure by benzimidazoles 3, yeast) |
|-------------|---|
| | oenermiraarotes o, reast, momorog (ru-gu.bedoorss. gb.Aro47472.1 |
| 201773_at | gb:NM_015339.1 /DEF-Homo sapiens activity-dependent neuroprotective protein (ADNP), mRNA. /FEA-mRNA /GEN-ADNP /PROD-activity-dependent neuroprotective protein /DB_XREF-gi:12229216 /UG-Hs.3657 activity-dependent neuroprotective pro- |
| 218751_s_at | rein /ru-gp:nr200000.1 gp:nr1 013330.1 gp:nr /ru-gp:nr200000.1 gp:nr1 013330.1 fp:nr /ru-gp:nr200000.1 gp:nr1 01301. |
| 201350_at | sapiens flotillin 2 (FLOT2), mRNA. /FEA=mRNA /GEN=FLOT2 /PROD=flotillin 2 2 FL=qb:NM 004475.1 qb:M60922.1 |
| 201098_at | 66.1 /DEF=Homo sapiens coatomer pro |
| | /GEN=COPB2 /PROD=coatomer protein complex, subunit beta 2 (betaprime) /DB_XREF=gi:4758031 /UG=Hs.75724 coatomer protein complex, subunit beta 2 (beta prime) /FL=gb:BC000326.1 qb:NM 004766.1 |
| 203732_at | 1 /DEF-Homo sapiens thyroid hormone receptor interactor 4 (TRIP4), mRNA. /FEA-mRNA /G |
| | /PROD-activating signal cointegrator 1 /DB_XREF=gi:7706430 /UG=Hs.116784 thyroid hormone receptor interactor 4 /FL-gb:AF168418.1 gb:NM 016213.1 |
| 202846_s_at | o sapiens phosphatidylinositol glycan, class C (PIGC), mRNA. /FEA=mRNA /GE |
| - | chass C /FLagb: D85418.1 gb: NM 002642.1 |
| 205936_s_at | gb:NM 002115.1 /DEF=Homo sapiens hexokinase 3 (white cell) (HK3), mRNA. /FEA=mRNA /GEN=HK3 /PROD=hexokinase 3 (white cell) /DB XREF=ci:4504394 /UG=Hs.159237 hexokinase 3 (white cell) /FL=ch:U51333 1 ch:NM 002115 1 |
| 201209_at | 004964.2 /DEF=Homo sapiens histone deacetylase 1 (HDAC1), mRNA. /FEA=mRNA /GEN=HDAC1 /PROD=histone deac |
| 217708 at | /UB |
| 25-0611 | Consensus includes grantzarzo (real-sar) DE Arter-grantzarzo (DE Arter-est: gatagous.xi /CLONE=indoe: 1690096 /UG=Hs.239720 CCR4-NOT transcription complex, subunit 2 /FIL-gb:ĀF113226.1 gb:AF180473.1 gb:NM 014515.1 |
| 209455_at | Consensus includes gb:BE963245 /FEA=EST /DB XREF=gi:11766663 /DB XREF=est:601656874R1 /CLONE=IMAGE:3865699 //OGHS.21229 f-box and WD-40 domain protein 1B /FL=qb;AF176022.1 qb:AB033281.1 |
| 212397_at | Consensus includes gb:AL137751.1 /DEF=Homo sapiens mRNA; cDNA DKFZp43410812 (from clone DKFZp43410812); partial cds. |
| | protein /DB_XREF=gi:680B387 /UG=Hs.263671 Homo sapiens mRNA; partial_cds |
| 218025_s_at | sapiens peroxisomal D3,D2-enoyl-CoA enoyl-CoA isomerase /DB_XREF=gi:5174 |
| | J02668.1 gb:AF069301.1 gb:AF153612.1 gb:NM 006117.1 gb:AF244138.1 |
| 203217_s_at | gb:NM 003896.1 /DEF=Homo sapiens sialyltransferase 9 (CMP-NeuAc:lactosylceramide alpha-2,3-sialyltransferase; GM3 syn- thase) (SIAT9), mRNA. /FEA=mRNA /GEN=SIAT9 /PROD=sialyltransferase 9 (CMP-NeuAc:lactosylceramidealpha-2,3- |
| : | <pre>/nthase) /DB XREF=g1:4506954 /UG=Hs.225939 sia 43 synthase) /FL=qb:AB018356.1 qb:NM 003896.1 db</pre> |
| 212812_at | 700633 /FER=EST /DB XREF=g1:4988533 /DB_XREF=e642 fis, clone HSI06970 |
| 207830_s_at | gb:NM_002713.1 /DEE=Homo sapiens protein phosphatase 1, regulatory (inhibitor) subunit 8 (PPPIR8), mRNA. /FEA=mRNA. //GRM=DPD188 /PRON=FROTE inhosphatase 1 regulatory (inhibitor) subunit 8 /DB VDEFECTION (INC-W 2005) |
| | hibitor) subunit 8 /FL=gb:NM 002713.1 gb:U14575 |
| 204061_at | gb:NM_005044.1 /DEF=Homo sapiens protein kinase, X-linked (PRKX), mRNA. /FEA=mRNA /GEN=PRKX /PROD=protein kinase, X- linked /DB XREF=gi:4826947 /UG=Hs.147996 protein kinase, X-linked /FL=gb:NM 005044.1 |

| 215633_x_at | ıt. |
|--------------------|--|
| 208660_at | |
| 218805_at | gb:NM 018384.1 /DEF=Homo sapiens hypothetical protein FLJ11296 (FLJ11296), mRNA. /FEA=mRNA /GEN=FLJ11296 /PROD=hypothetical protein FLJ11296 /DB XREF=qi:8922984 /UG=HS.26194 hypothetical protein FLJ11296 /FF=rh.NM 018384 1 |
| 217725_x_ | at db:NM_015640.1 /DEF=Homo sapiens PAI-1 mRNA-binding protein (PAI-RBP1), mRNA. /FEA=mRNA /GEN=PAI-RBP1 /PROD=F binding protein /DB XREF=q1:7661625 /UG=Hs.165998 PAI-1 mRNA-binding protein /FE=cb:BL080119.1 cb:NM_015640 |
| 200802_at | - |
| 213203_at | Consensus includes gb:A1633709 / small nuclear RNA activating com |
| 204640_s_a | gb:NM 003563.1 /DEF=Homo tein /DB XREF=gi:4507182 |
| 202324_s_at | gb:NM 022735.1 /DEF=Homo sapiens golgi resident protein GCP60 (GCP60), mRNA. /FEA=protein GCP60 /DB XREF=q1:12232390 /UG-Hs.6831 golgi resident protein GCP60 /FF=ab |
| 211661_x_at | gb:M80436.1 /DEF-Human platelet activating factor receptor mRNA, compl vating factor receptor /DB XREF=qi:189537 /FL=qb:M80436.1 |
| 208969_at | /FEA=mRNA /PROD=NA |
| * | iquinone) 1 alpha subcomplex, 9 |
| 202227_s_at | gb:NM 006696.1 /DEF=Homo /PROD=thyroid hormone rec |
| 221989_at | s includes gb:AW05 |
| 202545_at | |
| 202214_s_at | gb:NM 003588.1 /DEF=Homo sapiens cullin 4B (CUL4B), mRNA. /FEA=mRNA /GEN=CUL4B /PROD=cullin 4B /DB_XREF= //UG=HS.155976 cullin 4B /FL=gb:NM 003588.1 gb:AB014595.1 |
| 218366_x_at | et gb:NM 022734.1 /DEF=Homo sapiens hypothetical protein FLJ20859 (FLJ20859), mRNA. /FEA=mRNA /GEN=FLJ20859 (/ /PROD=hypothetical protein FLJ20859 /DB XREF=gi:12232388 /UG=Hs.6311 hypothetical protein FLJ20859 /FL=cp:NM 022734 1 |
| 218592 <u>s</u> at | gb:NM 017829.1 /DEF=Homo sapiens hypothetical protein FLJ20454 (FLJ20454), mRNA. //VPROD=hypothetical protein FLJ20454 /DB_XREF=gi:8923424 /UG=Hs.26890 hypothetical gb:NM 017829.1 |
| 209658_at | gb:AF164598.1 /DEF-Homo sapiens cell division control protein 16 (CDC16) mRNA, complete cds. /FEA=mRNA /GEN=CDC16 /PROD=cell division control protein 16 /DB_XREF=gi:5533374 /UG=Hs.1592 CDC16 (cell division cycle 16, S. cerevisiae, homolog) /FL=gb:AF164598.1 |
| 202217_at | gb:NM_004649.1 /DEF=Homo sapiens ES1 (zebrafish) protein, human homolog of (C210RF33), mRNA. /FEA=mRNA /GEN=C210RF33 / PROD=ES1 (zebrafish) protein, human homolog of /DB_XREF=gi:5031690 /UG=Hs.182423 ES1 (zebrafish) protein, human homolog of /EL=gb:BC002370.1 gb:BC003587.1 gb:D86061.1 gb:U53003.1 gb:NM_004649.1 |
| 54970_at | Homo sapiens cDNA, 3 end |

| 202441 at | Consensus includes gb:AL568449 /FEA=EST /DB_XREF=gi:12922799 /DB_XREF=est:AL568449 /CLONE=CSODE001YC12 (3 prime) |
|--------------|--|
| | /UG=HS.285818 similar to Caenorhabditis elegans protein C42Cl.9 /FL=gb:Atc0403.1 gb:NN UUB409.1 |
| 211383_s_at | apiens mkNa; cDNA DKE'264/41/2/2/ (IIOM CLOUE DKE'264/3/11/4/1/) COMPLECE CUE: hypothetical protein /DB XREF=gi:6807664 /UG=Hs.2727 KIA0982 protein /Fi- |
| 201903_at. | o sapiens ubiquinol-cytochrome c reductase core protein 1 (UCKCI), mkwA. ome c reductase core protein I /DB_XREF=gi:4507840 /UG=Hs.119251 ubiquinol- I /FiT=gb:L16842.1 gb:NM 003365.1 gb:D26485.1 |
| 201827_at | apiens PRO2451 mR sociated, actin d 1077.1 gb:AF113019 |
| 201583_s_at. | |
| 210825_s_at | gb.Ari30103.1 /DEF-Homo sapiens clone FLB2914 PRO0/20 mRNA, complete cds. /FL-BA-MRNA /FROU-ZO /DE_AREF-GILLIANSSOS /UG-Hs.160483 erythrocyte membrane protein band 7.2 (stomatin) /FL-BA-F130103.1 |
| 212696_s_at | 968633 /FEA=EST /L protein 4 |
| 218229_s_at | /DEF-Homo sapiens KIAA1513 protein (KIAA1513), 223835 /UG-Hs.284227 KIAA1513 protein /FI-gb:NM (|
| 210111_s_at | sapiens PNAS-138 mkNA, compiece cas. /rem-mkwa ns PNAS-138 mkNA, complete cds /rL-gb:AF277175. |
| 201268_at | <pre>gb:NM_002512.1 /DEF=Homo sapiens non-metastatic cells 2, protein (NM23B) expressed in (NME2), nuclear gene encoding mitochondrial protein, mRNA. /FEA=mRNA /GEN=NME2 /PROD=non-metastatic cells 2, protein (NM23B) expressed in /DB XREF=gi:4505408 /UG=Hs.275163 non-metastatic cells 2, protein (NM23B)</pre> |
| 201199_s_at | nacropain) 26S subunit, non ubunit,non-ATPase, 1 /DB_XR :D44466.1 gb:NM 002807.1 |
| 218242_s_at | sapiens hypothetical protein FLJ20039 (FLJ20039), mRNA. /FEA=mRNA /GEN=FLJ20039 in FLJ20039 /DB XREF=gi:8923045 /UG=Hs.267448 hypothetical protein FLJ20039 /FL=gb:NM |
| 201232_s_at | sapiens proteasome (prosome, macropain) 26S subunit, non-ATPase, 13 (PSML13), 1 some (prosome, macropain) 26S subunit, non-ATPase, 13 /DB XREF=gi:4506222 /UG=Hs subunit, non-ATPase, 13 /FL=gb:BC001100.1 gb:BC001147.1 gb:AB009398.1 |
| 220044_x_at | gb:NM_016424.1 /DEF=Homo sapiens cisplatin resistance-associated overexpressed protein (LUC/A), mRNA. /FEA=mRNA /GEN=LUC/A /PROD=cisplatin resistance-associated overexpressedprotein /DB_XREF=gi:7706534 /UG=Hs.3688 cisplatin resistance-associated overexpressed protein /FL=gb:NM 016424.1 |
| 210835_s_at | gb:AF222711.1 /DEF=Homo sapiens ribeye mRNA, complete cds. /FEA=mRNA /PROD=ribeye /DB_XREF=g1:12034652 /UG=HS.1/1391 C-terminal binding protein 2 /FI=gb:AF222711.1 gb:NM 022802.1 |
| 211271_x_at | act binding protein (neterogeneous nuclear ilar to polypyrimidine tract binding protein 50 polypyrimidine tract binding protein |
| 218998_at | 132.1 /DEF=Homo othetical prote 153417.1 gb:NM |
| 201765_s_at | Consensus includes gb:AL523158 /FEA=EST /DB_XREF=g1:12/86651 /DB_XREF=est:AL523158 /CLUONE=CSULCOLIEGO (5 PIIME) |

| | ase A (alpha polypeptide) /FL=gb:NM_000520.2 |
|-------------|---|
| 204837_at | 30178.1 /DEF=Homo sapiens mRNA; cDNA DKF2p434K171 (from clone DKrZp434K171 71 /PROD=hypothetical protein /DB_XREF=gi:5262652 /UG=Hs.27194 DKFZP434K17 |
| 201221_s_at | iomo sapiens small nuclear ribonucleoprotein 70kD polypepti lbonucleoprotein 70kD polypeptide(RNP antigen) /DB_XREF=gi: igen) /FL=gb:BC000342.1 gb:M22636.1 gb:NN 003089.1 |
| 209092_s_at | ns clone 016b03 My027 protein mRNA, complete cds. /FEA-mRNA /PROD-My027 protein /DB_XREF-g1:12001995 /FL-gb:AF061730.1 gb:AF151908.1 gb:NN 016080.1 |
| 202265_at | |
| 203721_s_at | gb:NM 016001.1 /DEF=Homo sapiens CGI-48 protein (LOC51096), mRNA. /FEA=mRNA /GEN=LOC51096 /PROD=CGI-48 protein /DB_XKEF=g1://U3/04 /UG-H5.6153 CGI-48 protein /FL=gb:AF151806.1 gb:NM 016001.1 |
| 202475_at | iens seven transmembrane domain protein (NIFIEL4), mRNA. /FEA-RNA /GEN-NIFIEL4 /PROD-seven transmembran 153781 /UG-Hs.9234 seven transmembrane domain protein /FI-59:BC001118.1 gb:N/ 006326. |
| 213165_at | 4 /FEA-EST /DB_XREF-gi:3280398 /DB_XREF-est:ov |
| 213198_at | JDEF-Homo sapiens mRNA; cDNA DKFZp434M245 cDNA DKFZp434M245 (from clone DKFZp434M245) |
| 212837_at | .1 /DEF-Human mRNA for KIAA0157 gene, partial cds. /FEA-mRNA /GEN-KIAA0157 /DB_KRE-g1:961445 |
| 206592_s_at | <pre>gb:NM 003938.1 /DEF=Homo sapiens adaptor-related protein complex 3, delta 1 subunit (AP3D1), mRNA. /FEA=mRNA /GEN=AP3D1 /PKON=adaptor- related protein complex 3, delta 1subunit /DB_XREF=gi:4501976 /UG=Hs.75056 adaptor-related protein complex 3, delta 1 subunit //FI=AP:NM 930.1 ob:NM 003938.1</pre> |
| 212694_s_at | 532.1 /DEF-Homo sapiens prop PA-CDS /GEN-PCCB /PROD-prop: Lase, beta polypeptide /FL-(|
| 211759_x_at | gb:BC005969.1 /DEE-Homo sapiens, clone MGC:14625, mRNA, complete cds. /FEA-mRNA /PROD-Unknown (protein for MGC:14625) /DB_XREE-g1:1334364 /FL-qb:BC005969.1 |
| 212714_at | Consensus includes gb:AL050205.1 /DEF-Homo sapiens mRNA; cDNA DKFZp586F1323 (from clone DKFZp586F1323). /FEA-mRNA /DB XREF-gi:4884444 /UG-Hs.26613 Homo sapiens mRNA; cDNA DKFZp586F1323 (from clone DKFZp586F1323) |
| 201528_at | XREF=est:602439 |
| 211795_s_at | <pre>gb:AF198052.1 /DEF-Homo sapiens EVH1 domain binding protein mRNA, complete cds. /FEA-CDS /PROD-EVH1 domain binding protein /DB XREF-qi:7416992 /UG-Hs.58435 FYN-binding protein (FXB-120130) /FL-gb:AF198052.1</pre> |
| 218633_x_at | pp:NM 018394.1 /DEE-Homo sapiens hypothetical protein FLJ11342 (FLJ11342), mRNA. /FEA-mRNA /GEN-FLJ11342 /FROD-hypothetical protein FLJ11342 /DB XREF-q1:8923000 /UG-Hs.266514 hypothetical protein FLJ11342 /FL-qb:NM 018394.1 |
| 200800_s_at | gb:NN 005345.3 /DEE=Homo sapiens heat shock 70kD protein 1A (HSPA1A), mRNA. /FEA=mRNA /GEN=HSPA1A /PROD=heat shock 70kD protein 1A /DB XREF=q1:5579469 /UG=Hs.8997 heat shock 70kD protein 1A /FL=gb:BC002453.1 gb:NN 005345.3 |
| 214719_at | =mRNA /DB_XREF=g1:10439638 /UG=Hs.1 |
| 213655_at | 1643 /FEA=EST /DB_XREF=gi:22376 protein, epsilon polypeptide |
| 212371_at | DKF2p586C1019) |
| 210166_at | gb:AF051151.1 /DEF=Homo sapiens Tollinterleukin-1 receptor-like protein 3 (TiL3) mkN4; complete cds. /red=mkN4 /GEN=11.3 /PROD=Tollinterleukin-1 /PROD=Tollinterleukin-1 receptor-like protein 3 /DB XREF=gi:3132525 /UG=H8.114408 toll-like receptor 5 /FL=gb:AF051151.1 |
| 217958_at | gb:NM 016146.1 /DEF=Homo sapiens PTD009 protein (PTD009), mRNA. /FEA=mRNA /GEN=PTD009 / PROD=PTD009 protein /DB_XREF=g1://D0000 //HG=N= 279501 prD009 |

| 219269_at | ens hypothetical protein FLJ21616 (FLJ21616), mRNA. /FEA-mRNA. /UG-Hs.14562 hypothetical protein FLJ21616 /FL-gb:NM 024567.1 |
|--------------------|---|
| 219574_at | gb:NM_017923.1 /DEE-Homo sapiens hypothetical protein FLJ20668 (FLJ20668), mRNA. /FEA-mRNA /GEN-FLJ20668 /PROD-hypothetical protein FLJ20668 /FL-gb:NM 017923.1 |
| 210183_x_at | gb:AF112222.1 /DEF-Homo sapiens nuclear protein SDK3 mRNA, complete cds. /FEA+mRNA /PROD-nuclear protein SDK3 /DB_XREF-gi:6563229 /UG-Hs.44499 pinin, desmosome associated protein /FL-gb:AF112222.1 |
| 211582_x_at | gb:AF000424.1 /DEF=Homo sapiens LST1 mRNA, cLST1C splice variant, complete cds. /FEA=mRNA /GEN=LST1 /DB_XREF=gi:2145063 /UG-Hs.88411 lymphocyte antigen 117 /FL=gb:AF000424.1 |
| 212557_at | Consensus includes gb:AB011148.1 /DEF-Homo sapiens mRNA for KIAA0576 protein, partial cds. /FEA-mRNA /GEN-KIAA0576 /PROD-KIAA0576 protein / DB XREF-g1:3043675 /UG-Hs.172329 KIAA0576 protein |
| 213387_at | Consensus includes gb:AB033066.1 /DEF-Homo sapiens mRNA for KIAA1240 protein, partial cds. /FEA-mRNA /GEN-KIAA1240 /PROD-KIAA1240 protein / DB XREF-g1:6330790 /UG-Hs.62576 KIAA1240 protein |
| 218143 <u>s</u> at | gb:NM 005697.2 /DEF-Homo sapiens secretory carrier membrane protein 2 (SCAMP2), mRNA. /FEA-mRNA /GEN-SCAMP2 /PROD-secretory carrier membra- ne protein 2 /DB_XREF-gi:5730030 /UG-Ms.238030 secretory carrier membrane protein 2 /FL-gb:BC001376.1 gb:BC004385.1 gb:AF005038.2 gb:NM 005697.2 |
| 211615_s_at | ne-rich protein mRNA, complete cds 2439.1 |
| 213359 at | Consensus includes gb:W74620 /FEA-ESI /DB XREF-gi:1384833 /DB XREF-est:zd77e04.s1 /CLONE-IMAGE:346686 /UG-Hs.303627 heterogeneous nuclear ribonucleoprotein D (AU-rich element RNA-binding protein 1, 37kD) |
| 217842_at | |
| 207170_s_at | |
| 214352_s_at | 99 /FEA=EST /DB_XREF=qi:11947594 /DB_XREF=est:602136427F1 /CLONE=INAG homolog |
| 218287_s_at | ens 2C, |
| 204661_at | |
| 205545_x_at | |
| 213750_at | Consensus includes gb:AA928506 /FEA-EST /DB_XREF=gi:3076797 /DB_XREF=est:om17g03.sl /CLONE=IMAGE:1541332 /UG=Hs.10762 ESTs |
| 221970_s_at | Consensus includes gb:AU158148 /FEA-EST /DB_XREF-g1:11019669 /DB_XREF-est:AU158148 /CLONE-PLACE1011221 /UG-Hs.321105 Homo sapiens cDNA: FLJ21737 fis, clone COLF3396 |
| 218684_at | gb:NM 018103.1 /DEF-Homo sapiens hypothetical protein FLJ10470 (FLJ10470), mRNA. /FEA-mRNA /GEN-FLJ10470 /PROD-hypothetical protein FLJ10470 /FEA-mRNA /GEN-FLJ10470 /PROD-hypothetical protein FLJ10470 /FEA-mRNA /GEN-FLJ10470 /PB XREF-g1:8922441./UG-Hs.44672 hypothetical protein FLJ10470 /FEA-gb:NM 018103.1 |
| 200628_s_at | gb:W61715.1 /DEF-Human tryptophanyl-tRNA synthetase (WRS) mRNA, complete cds. /FEA-mRNA /GEN-WARS /PROD-transfer RNA-Trp synthetase /DB XREF-g1:340367 /UG-Hs.82030 tryptophanyl-tRNA synthetase /EL-gb:M71804.1 gb:M61715.1 gb:NM 004184.2 |
| 203905_at | fic ribonuclease (deadenylation nuclease) (PARN /DB_XREF=gi:4505610 /UG=Hs.43445 poly(A)-specif |
| 204039_at | ens CCAATenhancer binding p XREF=gi:4757971 /UG=Hs.761 |
| 63009_at | Cluster Incl. AI188402:qd08e03.x1 Homo sapiens cDNA, 3 end /clone=IMAGE-1723132 /clone_end=3 /gb=AI188402 /g1=3739611 /ug=Hs.93391 //len=506 |
| 211922_s_at | gb:AY028632.1 /DEF-Homo sapiens catalase (CAT) mRNA, complete cds. /FEA-CDS /GEN-CAT /PROD-catalase /DB_XREF-gi:13562131 /FL-gb:AY028632.1 |
| 212861_at | Consensus includes gb:BF690150 /FEA-EST /DB_XREF=gi:11975558 /DB_XREF=est:602186478T1 /CLONE=IMAGE:4298635 /UG-Hs.19210 ESTs |

| 206715_at | gb:NM 012252.1 /DEF-Homo sapiens transcription factor EC (TEEC), mRNA. /FEA=mRNA./GEN=TFEC /PROD=transcription factor EC /PDB XREF-gi:6912701 /UG-Hs.113274 transcription factor EC /FL-ob:D43145.1 dh:NM 012252 1 |
|-------------------|--|
| 213123_at | |
| 204049_s_at | gb:NM 014721.1 /DEE-Homo sapiens KIPA0680 gene product (KIAA0680), mRNA. /FEA-mRNA /GEN-KIPA0680 /PROD-KIAA0680 gene product //OB XREF-gi:1662247 /UG-Hs.102411 KIAA0680 gene product /FI-cb:AB014580 1 ch.NA 014721 1 |
| . 201985_at | gb:NM 014846.1 /DEE=Homo sapiens KIAA0196 gene product (KIAA0196), mRNA. /FEA-mRNA /GEN=KIAA0196 /PROD-KIAA0196 gene product /FLS-ab:083780.1 ab:NM 014846.1 |
| 208773_s_at | gb:AL136943.1 /DEF-Homo sapiens mRNA; cDNA DKF2p58651024 (from clone DKF2p58661024); complete cds. /FEA-mRNA /GEN-DKF2p58661024 /PROD-hypothetical protein /DB XREF-g1:12053380 /UG-Hs.301226 KIAA1085 protein /FL-ch.Ali36443 1 |
| 219563_at | 9b:NM 024633.1 /DEF-Homo sapiens hypothetical protein FL/21276; mRNA. /FEA-MRNA /GEN=FL/21276 /PROD-hypothetical protein FL/21276 /FL-ob:NM 024633 1 FL/312786 /FL-ob:NM 024633 1 |
| 218501_at | gb:NM 019555.1 /DEE-Homo sapiens Rho guanine nucleotide exchange factor (GEF) 3 (ARHGEE3), mRNA. /FEA-mRNA /GEN-ARHGEF3 /PROD-Rho guanine nucleotide exchange factor (GEF) 3 /DB_XREF-g1:9506400 /UG-Hs.25951 Rho guanine nucleotide exchange factor (GEF) 3 /FL-gb:AF249744.1 gb:NM 019555.1 |
| 212833_at | Consensus includes gb: M74089.1 /DEE-Human TB1 gene mRNA, 3 end. /FEA-mRNA /GEN-TB1 /DB_XREE-g1:182400 /UG-Hs.75639 Human TB1 gene mRNA, 3 end |
| 209623 <u>a</u> t | Consensus includes gb:AW439494 /FEA-EST /DB XREF-egi:6974800 /DB XREF-est:xt19c01.x1 /CLONE-IMAGE:2779584 /UG-HS.167531 methylcrotonoyl- Coenzyme A carboxylase 2 (beta) /FL-gb:AB050049.1 gb:AF310971.1 gb:NF301000.1 gb:NM 022132.2 |
| 209969_s_at | Similar to signal transducer and activator of trans transducer and activator oftranscription 1, 91kD / ED /FL-qb:BC002704.1 |
| 219966_x_at | gb:NM 017869.1 /DEE-Homo sapiens BANP homolog, SMAR1 homolog (FLJ20538), mRNA. /FER-mRNA /GEN-FLJ20538 /FROD-BANP homolog, SMAR1 homolog / DB XREF-gi:8923506 /UG-Hs.194637 BANP homolog, SMAR1 homolog /FL-aps:NM 017869.1 |
| 213275_x_at | ím' |
| 210231_x_at | gb:D45198.1 /DEF=Human mRNA for template acyivating factor-I alpha, complete cds. /FEA-mRNA /GEN-set /PROD-template acyivating factor-I alpha /DB XREF-gi:971211 /UG-Hs.145279 SET translocation (m/eloid leukemia-associated) /Fi-ch-n45198 1 |
| 212474_at | Consensus includes gb:D87682.1 /DEF-Human mRNA for KIAA0241 gene, partial cds. /FEA-mRNA /GEN-KIAA0241 /DB_XREF-gi:1663699 /UG-Hs.150275 KIAA0241 protein |
| 208717_at | gb:BC001669.1 /DEF-Homo sapiens, Similar to oxidase (cytochrome c) assembly 1-like, clone MGC:2171, mRNA, complete cds. /FEA-mRNA / PROD-Similar to oxidase (cytochrome c) assembly1-like /DB_XREF-g1:12804516 /UG-Hs.151134 oxidase (cytochrome c) assembly 1-like /FET-gb:BC001669.1 gb:NM 005015.1 |
| 217527_s_at | Consensus includes gb:A1478300 /FEA=EST /DB XREF=g1:4371526 /DB XREF=est:tm39e01.x1 /CLONE=INAGE:2160504 /UG=Hs.192789 ESTs, Weakly similar to Alu6 HUMAN ALU SUBFAMILY SP SEQUENCE CONTANINATION WARNING ENTRY H. saciens |
| 220495_s_at | iens hypothe 6 /UG=Hs.106 |
| 200892_s_at | 12.2 |
| 201798_s_at | gb:NM 013451.1 /DEF=Homo sapiens fer-1 (C.elegans)-like 3 (myoferlin) (FERLUS), mRNA, /FER=mRNA /GEN=FERLUS /PROD=fer-1 (C.elegans)-like 3 (myoferlin) /DB XREF=gi:7305052 /UG=Hs.234680 fer-1 (C.elegans)-like 3 (myoferlin) /PI=ch:APS146 1 ch. NM 01345 1 ch. NM 0 |
| 202529_at | gb:NM 002766.1 /DEE=Homo sapiens phosphoribosyl pyrophosphate synthetase-associated protein 1 (PRPSAPI), mRNA. /FEA=mRNA /GEN=PRPSAPI / PROD=phosphoribosyl pyrophosphatesynthetase-associated protein 1 /DB_XREF=g1:4506130 /UG=Hs.77498 phosphoribosyl pyrophosphate synthetase-associated protein 1 /FL=qb:D61391.1 gb:NM 002766.1 |
| 208897_s_at | gb:BC003360.1 /DEF-Homo sapiens, DEADH (Asp-Glu-Ala-AspHis) box polypeptide 18 (Myc-regulated), clone NGC:5316, mRNA, complete cds. /FEA-mRNA /PROD-DEADH (Asp-Glu-Ala-AspHis) box polypeptide 18 (Myc-regulated) /DB_XREF-gi:13097182 /UG-Hs.100555 DEADH (Asp-Glu-Ala-AspHis) box polypeptide 18 (Myc-regulated) /FL-gb:BC001238.1 qb:BC003360.1 |
| 210296_s_at | <pre>gb:BC005375.1 /DEF=Homo sapiens, peroxisomal membrane protein 3 (35kD, Zellweger syndrome), clone MGC:12491, mRNA, complete cds. /FEA=mRNA /PROD=peroxisomal membrane protein 3 (35kD, Zellwegersyndrome) /DB_XREF=gi:13529226 /UG=Hs.180612 peroxisomal membrane protein 3 (35kD, Zellweger syndrome) /FI=gb:BC005375.1</pre> |
| 212460_at | Consensus includes gb: BE738425 /FEA-EST /DB_XREF-g1:10152417 /DB_XREF-est:60157244171 /CLONE-INAGE:3839147 /UG-Hs.241507 ribosomal protein S6 |

| 211800_s_at | ens deubiguitinating enzyme UnpES (UNP) mRNA, complete cds. /FEA-CDS /GEN-UNP /PR fic protease 4 (proto-oncogene) /FL-gb:AF01/306.1 |
|-------------|---|
| 202850_at | er |
| 212202_s_at | 72 /FER-EST /DB_XREF=g1:13455486 /DB_XREF-est:602542252F1 /CLONE-IMAGE:46 |
| 209337_at | iens lens epithelium-derived growth factor mRNA, complete cds. /FEA≒mRNA /PROD≒lens epithelium-derived /UG=Hs.82110 PC4 and SERS1 interacting protein 1 /FL-qb:NM 021144.1 gb:AF663020.1 |
| 207616_s_at | 35:NN 004180.1 /DEF-Homo sapiens TRAF family member-associated NFKB activator (TANK), mRNA. /FEA-mRNA /GEN-TANK /FROD-TRAF family member-associated NFKB activator /FL-qb:U63830.1 gb:NN 004180.1 |
| 204630_s_at | |
| 209422_at | 65 /DEF-Human DNA sequence from clone RP5-1121G12 on chromosome 20 Contains th carcinoma-associated antigen 58 (HCA58) and a putative novel protein containi B XREF-qi:9663113 /UG-Hs:112594 hypothetical protein DKF2p434F0272 /FL-gb:AV02 |
| 203658_at | /DEF=Homo sapi neacylcarnitine -gb:BC001689.1 |
| 212015_x_at | Consensus includes gb:BF690062 /FEA-EST /DB_XREF-gi:11975470 /DB_XREF-est:60218636611 /CLONE-IMAGE:4298440 /UG-Hs.172550 polypyrimidine tract binding protein (heterogeneous nuclear ribonucleoprotein I) |
| 222122_s_at | Consensus includes gb:BG403671 /FEA-EST /DB_XREF=g1:13297119 /DB_XREF=est:602419393F1 /CLONE=IMAGE:4526326 /UG-Hs.16411 hypothetical pro- cein LOC57187 |
| 202171_at | 275 /FEA-EST /DB_XREF-gi:11007796 /DB_XREF-est:AU146275 /CLONE-HEMBB1000004 /UG-Hs.6557 zinc 5.1 |
| 210460_s_at | IRNA for pUb-R5, complete cds. /FEA-mRNA /GEN-hpUb-R5 /PROD-F subunit, non-ATPase, 4 /FL-gb:AB033605.1. |
| 207551_s_at | |
| 201104_x_at | piens hypothetical protein (DJ328E19.C1.1), mRNA. /FEA=mRNA /GEN=DJ328E19.C1.1 /PROD=hypothetical prot 18329 hypothetical protein /FL=gb:NM 015383.1 |
| 200812_at | oiens chaperonin containing REF=gi:5453606 /UG=Hs.108809 |
| 203494_s_at | =Homo sapiens KIAA0092 gene product (KIAA0092), mRNA. /FEA=mRNA /GEN=KIAA0092 /PRC /UG-Hs.151791 KIAA0092 gene product /FL=gb:D42054.1 gb:NM 014679.1 |
| 202127_at | |
| 212532_s_at | .64 /FEA=EST /DB_XREF=g1:8007617 /DB_XREF=egt:ho62c10.x1 /CLONE=INAGE:3041970 /UG=Hs.15 |
| 218357_s_at | gb:NN 012459.1 /DEF-Homo sapiens translocase of inner mitochondrial membrane 8 (yeast) homolog B (TIMMBB), mRNA. /FEA-mRNA /GEN-TIMMBB / PROD-translocase of inner mitochondrial membrane 8(yeast) homolog B /DB_XREF-g1:6912711 /UG-Hs.279915 translocase of inner mitochondrial. |
| | 150087.1 gb: AF152350.1 gb: NM 012459.1 |
| 208847_s_at | gb:N29872.1 /DEF-Human alcohol dehydrogenase class III (ADH5) mRNA, complete cds. /FEA-mRNA /GEN-ADH5 /DB_XREF-gi:178131 /UG-Hs.78989 alco-hol dehydrogenase 5. (class III), chi polypeptide /FL-gb:NN 000671.2 gb:M29872.1 gb:M30471.1 |
| 207730_x_at | pier 9 /(|
| 212896_at | 1.2 /DEF=Homo sapiens mRNA for KIAA0052 protein, partia 18608 KIAA0052 protein |
| 203156_at | oiens A-kinase anchoring protein 220 (LOC51707), mRNA. /FEA-mRNP -Hs.232076 A kinase (PRKA) anchor protein 11 /EL-gb:AF176555.1 g |
| 220066_at | gb:NM 022162.1 /DEF=Homo sapiens NOD2 protein (NOD2), mRNA. /EEA=mRNA /GEN=NOD2 /PROD=NOD2 protein /DB_XREF=gi:11545911 /UG=Hs.135201 NOD2 |

| | protein /FL-gb:AF178930.1 gb:NM_022162.1 |
|--------------------|---|
| 218583_s_at. | 95:NM 020640.1 /DEF=Homo sapiens RP42 homolog (RP42), mRNA. /FEA-mRNA /GEN-RP42 /PROD-RP42 homolog /DB_XREF-gi:10190677 /UG-Hs.104613 RP42 homolog /FL-gb:NM 020640.1 gb:AF292100.2 |
| 221689_s_at | gb:AB035745.1 /DEF-Homo sapiens mRNA for DSCR5b, complete cds. /FEA-mRNA /GEN-DSCR5b /PROD-DSCR5b /DB_XREF-gi:7798596 /UG-Hs.66493 Down syndrome critical region gene 5 /FL-gb:AB035745.1 gb:AB037163.1 dp:AF237812.1 |
| 205412_at | ens acetyl-Coenzyme A acetyltransf /FEA=mRNA /GEN=ACAT1 /PROD=acetyl ferase 1 (acetoacetyl Coenzyme A t |
| 202542_s_at | ens small inducible cytokine subfamily E, ble cytokine subfamily E, member 1 /DB XRE, activating) /Fl=qb:NM 004757.1 qb:U10117.1 |
| 202521_at | ens CCCTC-binding factor (linc finger protein) (CTCE), mRNA. /FEA-mRNA /GEN-CTCE /PROD- F-q1:5729789 /UG-Hs.57419 CCCTC-binding factor (linc finger protein) (refer to Acted) |
| 204391_x_at | gb:NM_015905.1 /DEF=Homo sapiens transcriptional intermediary factor 1 (TIF1), mRNA. /FEA-mRNA /GEN-TIF1 /PROD-transcriptional intermediary factor 1 /FL=gb:AF009353.1 gb:AF119042.1 gb:NM_003852.1 gb:NM_003852.1 |
| 214733_s_at | Consensus includes gb:AL031427 /DEE=Human DNA sequence from clone 167A19 on chromosome 1p32.1-33. Contains three genes for novel proteins, the DIO1 gene for type I iodothyronine deiodinase (EC 3.8.1.4, TXDI1, ITDI1) and an HNRNP A3 (Heterogenous Nuclear Ribonucleoprotein A3, FERA-MRNA 6 /DB XREE=g1:4835258 /UG-HS.11923 hypothetical protein |
| 217864_s_at | gb:NN 016166.1 /DEF=Homo sapiens DEADH (ASp-Glu-Ala-AspHis) box binding protein 1 (DDXBP1), mRNA. /FEA=mRNA /GEN=DDXBP1 /PROD=DEADH (Asp-Glu-Ala-AspHis) box binding protein 1 /FL=gb:AF077951.1 gb:AF167160.1 gb:NN 016166.1 |
| 212904_at | Consensus includes gb:AB033011.1 /DEF-Homo sapiens mRNA for KIAA1185 protein, partial cds. /FEA-mRNA /GEN-KIAA1185 /PROD-KIAA1185 protein / DB XREF-qi:6330301 /UG-Hs.268488 KIAA1185 protein |
| 202126_at | Consensus includes gb:AA156948 /FEA=EST /DB XREF=gi:1728563 /DB XREF=est:2119f02.s1 /CLONE=IMAGE:502395 /UG-Ms.198891 serinethreonine- protein kinase PRP4 homolog /FL=ab:U48736.1 ab:WM 003913.1 |
| 203484_at | piens Sec61 gamma NM 014302.1 |
| 203345_s_at | Consensus includes gb:AI566096 /FEA-EST /DB_XREF-gi:4524548 /DB_XREF-est:tn53d02.x1 /CLONE-INAGE:2172099 /UG-Hs.31016 putative DNA binding pictein /FL-gb:AF072814.1 gb:NM 007358.1 |
| 213 2 38_at | Consensus includes gb:AI478147 /FEA-EST /DB_XREF-g1:4371373 /DB_XREF-est:tm34f06.x1 /CLONE-INAGE:2160035 /UG-Hs.173540 ATPase, Class V, type 10D |
| 202680_at | gb:NM 002095.1 /DEF-Homo sapiens general transcription factor IIE, polypeptide 2 (beta subunit, 34kD) (GTEZEZ), mRNA. /FEA-mRNA. /GEN-GTEZEZ /PROD-general transcription factor IIE, polypeptide 2 (beta subunit, 34kD) /DB_XREF-g1:4504194 /UG-Hs.77100 general transcrip- tion factor IEE. polybeptide 2 (beta subunit, 34kD) /FI-ch-NA 002065 1 |
| 218117_at | /DEF-Homo sapiens rir 6:1 gb:AF140598.1 gb: |
| 218768_at | s nuclear pore |
| 202271_at | Consensus includes gb:AB007952.1 /DEF-Homo saplens mRNA for KIAA0483 protein, partial cds. /FEA-mRNA /GEN-KIAA0483 /PROD-KIAA0483 protein / DB XREF-gi:3413925 /UG-Hs.64691 KIAA0483 protein /FL-gb:NM 015176.1 |
| σ. | gb:NM 022750.1 /DEF-Homo sapiens hypothetical protein FLJ226931 (FLJ226931, mRNA. /FEA-mRNA /GEN-FLJ22693 /PROD-hypothetical protein FLJ22693 /DB XREF-g1:12232412 /UG-Hs.12646 hypothetical protein FLJ22693 /FL-ab:AL136766.1 qb:NM 022750.1 |
| 203146_s_at | gb:NM 001470.1 /DEF-Homo sapiens gamma-aminobutyric acid (GABA) B receptor, 1 (GABBR1), transcript variant 1, mRNA. /FEA-mRNA /GEN-GABBR1 / PROD-gamma-aminobutyric acid (GABA) B receptor 1, isoform a precursor /DB_XREF-gi:10835014 /UG-Hs.167017 gamma-aminobutyric acid (GABA) B receptor, 1 /FL-gb:NM 001470.1 gb:AF301005.1 gb:AF509148.1 |
| 218140_x_at | |
| 40420_at | Cluster Incl. AB015718:Homo sapiens lok mRNA for protein kinase, complete cds /cds=(50,2956) /gb=AB015718 /gi=4001687 /ug=Hs.16134 //en=4221 |

| <u>12.7</u> | at | 9 |
|-------------|-------------|---|
| | 48580_at | Cluster Incl. U55777:HSU55777 Homo sapiens cDNA /clone-37698 /gb-U55777 /gi-1354513 /ug-Hs.180933 /len-1593 |
| | | ens tetratricopeptide repeat domain 2 (TTC2), mRNA. /FER-mRNA /GEN-TTC2 /PROD-tetratricop 5542 DnaJ (Hsp40) homolog, subfamily C. member 7 /FL-gb:U46571.1 gb:NM 003315.1 |
| | 211727_s_at | ns, COX11 (yeast) homolog, cytochrome c oxidase assembly protein, clone MGC:14469, mRNN, complete homolog, cytochrome c oxidaseassembly protein /DB XREF=gi:13543474 /FL=gb:BC005895.1 |
| • | 202125_s_at | ens amyotrophic lateral sclerosis 2 (juvenile) chromosome region, candidate 3 (ALS2CR3), mRNA. /FEA=mF c lateral sclerosis 2 (juvenile)chromosome region, candidate 3 /DB_XREF=gi:13027379 /UG=Hs.154248 amyot s) chromosome region, candidate 3 /FL=gb:AB038951.1 gb:NM 015049.1 |
| · | 209451_at | gb:US9863.1 /DEF-Human TRAF-interacting protein I-TRAF mRNA, complete cds. /FEA=mRNA /PROD=I-TRAF /DB_XREF=gi:1518017 /UG=Hs.146847 TRAF family member-associated NFKB activator /FL=gb:US9863.1 |
| | 209440_at | ıns, phosphoribosyl pyrophosphate synthetase 1, clone MGC:2256, mRNA, complete cds. /FEA-mRNA sphate synthetase 1 /DB_XREE=g1:12804406 /UG-Hs.56 phosphoribosyl pyrophosphate synthetase 1 // |
| | 218189_s_at | ens N-acetylneuraminic acid phosphate synthase; stalic acid syntha phosphate synthäse:/DB_XREF-gi:12056472 /UG-Hs.274424 N-acetylneur b:BC000008:1 gb:AF257466.1 |
| | 200947_s_at | ens glutamate dehydrogenase 1 (GLUD1), mRNA. /FEA=mRNA /GEN=GLUD1 /PROD=glutamate dehydrogenase 1 1508 glutamate dehydrogenase 1 /FL=gb:J03248.1 gb:N37154.1 gb:M20867.1 gb:NM 005271.1 |
| | 202798_at | ens SEC24 (S. cerevisiae) related gene family, member B (SEC24B), mRNA. /FEA-mRNA /GEN-SEC24B /PROD- ly,member B /DB_XREF-gi:5454045 /UG-Hs.7239 SEC24 (S. cerevisiae) related gene family, member B |
| | 205323_s_at | iens metal-regulatory transcription factor 1 (MTF1), mRNA. /FEA-mRNA /GEN-WTF1 /PROD-metal-regula gi:5174588 /UG-Hs.211581 metal-regulatory transcription factor 1 /FL-gb:NM 005955.1 |
| | 212622_at | /FEA-EST /DB_XREF-g1:1212589 /DB_XREF-est:yz30c06.sl /CLONE-IMAGE:284554 /UG-Hs.174905 KIAP |
| | 212033_at | 17 /FER-EST /DB_XREF=g1:10809003 /DB_XREF=est:7j75a05.x1 /CLONE-IMAGE:3392240 /UG-Hs.180789 S16 |
| | 218919_at | /GEN=FLJ14007 / |
| | 209271_at | |
| | 213322_at | Consensus includes gb:AL031778 /DEF-Human DNA sequence from clone 34B21 on chromosome 6p12.1-21.1. Contains part of a gene for a novel |
| | | bert of Tight Junction Protein 201: [TJP1] and UNC5 Homologs, the gene for a novel B2RP (peripheral benzodiazap (REF-g1:1153958 /UG-Hs.183056 Human DNA sequence for a conclusion of themseome 6p12.1-21.1. Contains part of a |
| | | נס part סו ווקור טעוכרוטו גיסרפוו נטו (נטגו) מוט טענט מטווטוטעט, רוופ קפופ זטו מ ווטיפו |
| | 221895_at | 34 /FEB=EST /DB_XREF=g1:7039290 /DB_XREF=est:hc78g04.x1 /CLONE=IMAGE:2898870 /UG=Hs.65400 |
| | 217760_at | 80 /FER=EST /DB_XREF=g1:1757929 /DB_XREF=est:zp32a10.s1 /CLONE=INAGE:611130 /UG=Hs.14512 DIPB protei |
| | 201448_at | |
| | 218465_at | |
| | 212880_at | /DEF=Homo sapiens mRNA for KIAA0541 protein, partial cds. /FEA=mRNA /GEN=KIAA W repeat domain 7 |
| | 218142_s_at | ens protein x 0001 (LOC51185), mRNA. /FEA-mRNA /GEN-LOC51185 /PROD-protein x 0001 /DB_XRE 'FI-gb:NM 016302.1 gb:AF117230.1 |
| | 202078_at | <pre>gb:NM 003653.1 /DEF=Homo sapiens COP9 (constitutive photomorphogenic, Arabidopsis, homolog) subunit 3 (COPS3), mRNA. /FEA=mRNA /GEN=COPS3 / /PROD=COP9 (constitutive photomorphogenic, Arabidopsis, homolog) subunit 3 /DB XREF=gi:4502974 /UG=Hs.6076 COP9 (constitutive photomorpho-</pre> |

| | genic, Arabidopsis, homolog) subunit 3 /FL=gb:BC001891.1 gb:AF031647.1 gb:NM_003653.1 gb:AF098109.1 |
|-------------|--|
| 219069_at | gb:NM 017704.1 /DEF=Homo sapiens hypothetical protein FLJ20189 (FLJ20189), mRNA. /FEA=mRNA /GEN=FLJ20189 /PROD-hypothetical protein FLJ20189 /FL=qb:NM 017704.1 |
| 218333_at | gb:NM 016041.1 /DEE=Homo sapiens CGI-101 protein (LOC51009), mRNA. /FEA=mRNA /GEN=LOC51009 /PROD=CGI-101 protein /DE_XREF=g1:7705603 /VG=HS.286131 CGI-101 protein /FL=gb:AF132289.1 gb:NR 016041.1 gb:NR 016041.1 gb:RF242523.1 |
| 213082_s_at | Consensus includes 9b:AJ005866.1 /DEF=Homo sapiens mRNA for putative Sqv-7-like protein, partial. /FEA=mRNA /PROD=Sqv-7-like protein / DB XREF=gi:4008516 /UG-Hs.90078 nucleotide-sugar transporter similar to C. elegans sqv-7 |
| 221452_s_at | gb:RM 030969.1 /DEF=Homo sapiens hypothetical protein MGC1223 (MGC1223), mRNA. /FEA=CDS /GEN=MGC1223 /PROD=hypothetical protein MGC1223 / DB XREF=g1:13624338 /FL-gb:NM 030969.1 |
| 213733_at | Consensus includes gb:BF740152 /FEA-EST /DB_XREF-gi:12066828 /DB_XREF-est:7nl3g08.xl /CLONE=IMAGE:3564495 /UG-Hs.121555 myosin IE |
| 201225_s_at | gb:NM_005839.1 /DEE=Homo sapiens SerArg-related nuclear matrix protein (plenty of prolines 101-like) (SRN160), mRNA. /FEA-mRNA /GEN-SRN160 /PROD=SerArg-related nuclear matrix protein (plenty of prolines 101-like) /DB_XREF=gi:5032118 /UG=Hs.18192 SerArg-related nuclear matrix brotein (plenty of prolines 101-like) /FL-qb:AF048977.1 qb:NM_005839.1 |
| 219392_x_at | piens hypotl 1 /UG-Hs.27 |
| 219212_at | gb:NM 016299.1 /DEE-Homo sapiens heat shock protein hsp70-related protein (LOC51182), mRNA. /FEA-mRNA /GEN-LOC51182 /FROD-heat shock protein hsp70-related protein /DB_XREF-gi:10047093 /UG-Hs.44581 heat shock protein hsp70-related protein /EL-gb:NM_016299.1 gb:AF112210.1 gb:AF143723.1 |
| 217907_at | gb:NN 014161.1 /DEE=Homo sapiens HSPC071 protein (HSPC071), mRNA. /FEA-mRNA /GEN-HSPC071 /PROD-HSPC071 protein /DB_XREF-gi:7661777 /UG-HS.23038 HSPC071 protein /FL-gb:AL136633.1 gb:BC001623.1 gb:RF161556.1 gb:NN 014161.1 |
| 200727_s_at | Consensus includes gb:NA699583 /FEA-EST /DB XREF-g1:2103730 /DB XREF-est:zi42g07.sl /CLONE-IMAGE:433500 /UG-Hs.42915 ARP2 (actin-related protein 2, yeast) homolog /FI-gb:AF006082.1 gb:NF 005722.1 |
| 209068_at | U-rich element heterogeneous r |
| 46256_at | 05.s1 Homo sapiens cDNR, 3 end /clone=INAGE-979184 /clone_end=3 /gb=AA522 |
| 218068_s_at | gb:NN 024836.1 /DEE-Homo sapiens hypothetical protein FLJ22301 (FLJ22301), mRNN. /FEA-mRNA /GEN-FLJ22301 /PROD-hypothetical protein FLJ22301 /DB XREF-gi:13376246 /UG-Hs.181406 hypothetical protein FLJ22301 /FL-qb:NM 024836.1 |
| 208843_s_at | one NGC:1233, mRNA, complete cds. /FEA=n =gb:BC001408:1 |
| 202869_at | s 2,5-oligoadenylate synthetase 1 (10-46 kD) se 1, isoform E18 /DB_XREF=g1:8051620 /UG=Hs |
| 213546_at | Consensus includes gb:ALO50378,1 /DEF-Homo sapiens mRNA; cDNA DKF2p58611420 (from clone DKF2p58611420); partial cds. /FEA-mRNA /GEN-DKF2p58611420 /PROD-hypothetical protein /DB_XREF-g1:4914581 /UG-Hs.112423 Homo sapiens mRNA; cDNA DKF2p58611420 (from clone DKF2p58611420); partial cds |
| 214789_x_at | Consensus includes gb:AA524274 /FEA-EST /DB_XREF=g1:2265202 /DB_XREF=est:ng34dDB.sl /CLONE=IMAGE:936687 /UG-Hs.155160 Splicing factor, arginineserine-rich, 46kD |
| 213649_at | Consensus includes gb: PA524053 / FEA-EST / DB_XREF=gi:2264981 / DB_XREF=est:ng33b07.sl / CLONE-IMAGE:936565 / UG-Hs.184167 splicing factor, archineserine-rich 7 (35kD) |
| 212251_at | Consensus includes gb: A1972475 / FEA-EST / DB XREF-g1:5769391 / DB XREF-est:wr40b09.xl /CLONE-IMAGE:2490137 / UG-Hs.243901 Homo sapiens CDNA FL320738 fis, clone HEP08257 |
| 209166_s_at | gb:U68567.1 /DEF=Human lysosomal acid alpha-mannosidase mRNA, complete cds. /FEA=mRNA /PROD=lysosomal acid alpha-mannosidase // DB XREF=g1:1658373 /UG-Hs.279854 mannosidase, alpha, class 2B, member 1 /FL-gb:NM_000528.1 gb:U05572.1 gb:U65572.1 gb:U68567.1 gb:U60266.1 |
| 200683_s_at | 89 /FEA=EST /DB_XREF=g1:11768267 /DB_XR =gb:NM 003347.1 |
| 219913_s_at | gb:NM 016652.2 /DEF-Homo sapiens CGI-201 protein (LOC51340), mRNA. /FEA-mRNA /GEN-LOC51340 /PROD-CGI-201 protein /DB_XREF-gi:11072090 /UG-HS.268281 crooked neck protein (crn) /FL-gb:AF255443.2 gb:NM 016652.2 gb:AF318302.1 |
| 218104_at | gb:NN 017146.1 /DEF=Homo sapiens hypothetical protein FLJ20287 (FLJ20287), mRNA. /FEA=mRNA /GEN-FLJ20287 /FROD-hypothetical protein FLJ20287 /FLJ20287 /DB XREF=gi:8923268 /UG=Hs.26369 hypothetical protein FLJ20287 /FL-gb:NM 017746.1 |

| 212345_s_at | Consensus includes gb:BE675139 /FEA-EST /DB_XREF-gi:10035680 /DB_XREF-est:7f03b12.x1 /CLONE-IMAGE:3293567 /UG-Hs.13659 hypothetical protein pKFzp586F2423 |
|--------------|--|
| 213982_s_at | Consensus includes gb:BG107203 /FEA-EST /DB_XREF-gi:12601049 /DB_XREF-est:602290933F1 /CLONE=IMAGE:4385577 /UG-Hs.242271 KIAA0471 gene product |
| 202519_at | gb:NM_014938.1 /DEF-Homo sapiens KIAA0867 protein (MONDOA), mRNA. /FEA-mRNA /GEN-MONDOA /PROD-MondoA protein /DB_XREF-gi:7662347 /UG-Hs.52081 KIAA0867 protein /FL-gb:AB020674.1 gb:NM 014938.1 |
| 202892_at | gb:NM_004661.1 /DEF-Homo sapiens CDC23 (cell division cycle 23, yeast, homolog) (CDC23), mRNA. /FEA-mRNA /GEN-CDC23 /PROD-cell division cycle 23, yeast homolog; CDC23 /DB_XREF-gi:4757947 /UG-Rs.153546 CDC23 (cell division cycle 23, yeast, homolog) /FL-gb:AF053977.1 gb:AB011472.1 gb:AF01111 gb:AF191341.1 |
| 213876_x_at. | Consensus includes gb:AW089584 /FEA=EST /DB XREF=g1:6046928 /DB XREF=est:xd20f04.xl /CLONE=IMAGE:2594335 /UG-Hs.171909 U2 small nuclear ribonucleoprotein auxiliary factor, small subunit 2 |
| 202054_s_at | gb:NM 000382.1 /DEF-Homo sapiens aldehyde dehydrogenase 3 family, member A2 (ALDH3A2), mRNA, /FEA-mRNA /GEN-ALDH3A2 /PROD-aldehyde dehy-drogenase 3A2 /DB XREF-gi:4557302 /UG-Ms.159608 aldehydrogenase 3 family, member A2 /FL-gb:L47162.1 gb:U46689.1 gb:NM 000382.1 |
| 217776_at | ns androgen-regulated short-chain dehydrogenasereductase 1 (ARSDR1) mRNA, complete cds. / t-chaindehydrogenasereductase 1 /DB_XREF=g1:9622123 /UG-Hs.179817 CGI-82 protein /FL=gb:BC gb:AF167438.1 |
| 208758_at | gb:D89976.1 /DEE-Momo sapiens mRNN for 5-aminoimidazole-4-carboxamide ribonucleotide transformylase, complete cds. /FEA-mRNN / PROD-5-aminoimidazole-4-carboxamide ribonucleotidetransformylase /DB XREF-g1:2317691 /UG-Hs.90280 5-aminoimidazole-4-carboxamide ribonucleotide formyltransferaseINP cyclohydrolase /FL-gb:U37436.1 gb:D82348.1 gb:D89976.1 gb:NN 004044.1 |
| 212749_s_at | Consensus includes gb:AI096477 /FEA-EST /DB_XREF-gi:3445971 /JDB_XREF-est:qa03c06.xl /CLONE-IMAGE:1685674 /UG-Hs.48297 DKF2P586C1620 protein |
| 200890_s_at | Consensus includes gb:AW006345 /FEA-EST /DB_XREF-gi:5855123 /DB_XREF-est:wt04d05.xl /CLONE-IMAGE:2506473 /UG-Hs.250773 signal sequence receptor, |
| | |
| 211675_s_at | gb:AF054589.1 / DEF-Homo sapiens HIC protein isoform p40 and HIC protein isoform p32 mRNAs, complete cds. /FEA-mRNA /PROD-HIC protein iso- form p32; HIC protein isoform p40 /DB XREF-g1:3426297 /FL-gb:AF054589.1 |
| 206792_x_at | gb:NM 000923.1 / DEF-Homo sapiens phosphodiesterase 4C, cAMP-specific (dunce (Drosophila)-homolog phosphodiesterase E1) (PDE4C), mRNM. - FER-mRNM /GEN-PD61C / PROD-phosphodiesterase 4C, cAMP-specific (dunce(Drosophila)-homolog phosphodiesterase E1) / DB_XREF-g1:4505664 |
| , A.S. | ila)-homolog phosphodiesterase E1) |
| 218718_at | gb:NM 016205.1 /DEF=Homo sapiens platelet derived growth factor C (PDGEC), mRNA. /FEA=mRNA /GEN=PDGFC /PROD-secretory growth factor-like protein fallotein /DB_XREF=gi:9994186 /UG=Hs.43080 platelet derived growth factor C /FL=gb:AF091434.1 gb:AF244813.1 gb:AB033831.1 gb:NH 016205.1 |
| 201742_x_at | gb:NM 006924.1 /DEF-Homo sapiens splicing factor, arginineserine-rich 1 (splicing factor 2, alternate splicing factor) (SFRSI), mRNA. /FEA-mRNA /GEN-SFRS1 /PROD-splicing factor, arginineserine-rich 1(splicing factor 2, alternate splicing factor) /DB XREF-gi:5902075 //UG-Hs.73737 splicing factor, arginineserine-rich 1 (splicing factor 2, alternate splicing factor) /FL-db:M69040.1 db:NN 006044 |
| 218622_at | 1 |
| 212176_at | Consensus includes gp:AA902326 /FEA-EST /DB_XREF-g1:3037233 /DB_XREF-est:0192b01.sl /CLONE-IMAGE:1521385 /UG-Hs.18368 DKF2P564B0769 protein |
| 218970_s_at | gb:NM 015960.1 /DEF-Homo sapiens CGI-32 protein (LOCS1076), mRNA. /FEA-mRNA /GEN-LOCS1076 /PROD-CGI-32 protein /DB_XREF-g1:7705727 / UG-Hs.16606 CGI-32 protein /FL-gb:AF132966.1 gb:NM 015960.1 |
| 201696_at | P 81 |
| 222103_at | ᇬᆌ |
| 204036_at | 5 /FEA=EST /DB_XREF=gi:6656365 /DB_XREF d G-protein-coupled receptor, 2 /FL=gb: |
| 218283_at | gb:NN 016305.1 /DEF=Homo sapiens kiaa-iso protein (LOC51188), mRNA. /FEA=mRNA /GEN=LOC51188 /PROD=kiaa-iso protein /DB_XREF=gi:10047103 /UG=Hs.9774 kiaa-iso protein /EL=gb:NN 016305.1 gb:AF201950.1 |
| 209654_at | gb:BC004902.1 /DEF-Homo sapiens, clone MGC:4271, mRNA, complete cds. /FEA-mRNA /PROD-Unknown (protein for MGC:4271) /DB_XREF-gi:13436178 /UC-Hs.5070 KIAA0947 protein /EL-gb:BC004902.1 |

| 209083_at | gb:U34690.1 /DEF=Human coronin-like protein (HCORO1) mRNA, complete cds. /FEA-mRNA /GEN-HCORO1 /PROD-coronin-like protein //DB XREF=q1:1002922 /UG-Hs.109606 coronin. actin-binding protein 12 /FI=nh:134600 1 ch:NM 003024 1 ch:04.02 |
|--------------|--|
| 221873_at | 15 /FEA-EST /DB_XREF=gi:6301048 /DB_XREF=est:au72d09.x1 /CLOND=IRAGE:278 |
| 213573_at | Consensus includes gb:AA861608 /FEA-EST /DB_XREF-gi:2953748 /DB_XREF-est:ak34e01.s1 /CLONE=IMAGE:1407864 /UG=Hs.180446 karyopherin (im-portin) beta 1 |
| 209580_s_at | gb:AF114784.1 /DEF=Homo sapiens methyl-CpG binding endonuclease (MED1) mRNA, complete cds. /FER=mRNA /GEN=MED1 /PROD=methyl-CpG binding endonuclease /DB XREF=q1:4539758 /UG=Hs.35947 methyl-CpG binding domain protein 4 /Fi=rh-BF072750 1 ch-Na. A02005 1 - 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 |
| 219097_x_at | 1 / DEF-Homo sapiens hypothetical protein MGC2747 (MGC2747), mRNA. /FE |
| 206513_at | iens absent in mela noma 2 /FL=qb:AF02 |
| 209268_at | ens vacuolar protein sorting 49 KREF-qi:9294732 /UG-Hs.6650 vac |
| 219329_s_at | sapiens apoptosis related protein APR-3 (APR-3), mRNA. /FEA-mRNA /GEN-APR-3 /PROD-apoptosis re |
| | optosis related protein |
| 202051_s_at | gp:NM U05095.1 /DEE=Homo sapiens zinc finger protein 262 (zNF262), mRNA. /FEA=mRNA /GEN=ZNF262 /PROD=zinc finger protein 262 //DB XRFE-gi:s927068 / STE-FE - STERMEN STATE STA |
| 200749_at | Consensus includes gb:BF112006 /FEA=EST /DB XREF=q1:10941619 /DB XREF=est:7137e05.x1 /CLONF∈TMAGF:3527665 /HG=Hc 10842 bhW moments |
| | Oncogene family / [FL=qb:BC004272.1 qb:M31469.1 qb:AF052578.1 ob:AF054183.1 nh:NM ON6308.2 |
| 213373_s_at. | ncludes gb:BF439983 /FEA-EST /DB_XREF⊸gi:11452500 /DB_XREF=e |
| • | • |
| 212673_at | |
| 201771 at | iens secretoru catrior membrana mestais 3 (conuncia - mus. /mus.) |
| | 2076 |
| 209607_x_at | gb:U08032.1 /DEF=Numan thermolabile (monoamine, M form) phenol sulfotransferase (STM) mRNA, complete cds. /FEA=mRNA /GEN=STM /PROD=thermolabile (monoamine, M form) phenolsulfotransferase /DB XREF=gi:468256 /UG=Hs.274614 sulfotransferase family, cytosolic, 1A, phenol-preferring, member 3 /FE=ab:MM 003166.1 ab:119956.1 ab:114199 1 ab:114199 |
| 217317_s_at | 9b:AB002391.2 /DEF-Homo sapiens mRNA for KIAA0393 pv /UG-Hs.266933 hect domain and RLD 2 |
| 202272_s_at | KIAA0483 protein (K) |
| 210629 x at | gb:AF000425.1 /DEF=Homo sapiens LST1 mRNA, cLST1A splice variant, complete cds. /FEA-mRNA /GEN=LST1 /DB_XREF=g1:2145065 /UG=Hs.88411 lymphocyte antiqen 117 /FL=ab:AF000425.1 |
| 205329_s_at | gb:NM 003794.1 /DEF=Homo sapiens sorting nexin 4 (SNX4), mRNA. /FEA-mRNA /GEN=SNX4 /PROD-sorting nexin 4 /DB_XREF=g1:4507144 /UG-Hs.267812 sorting nexin 4 /FL-qb:AF130078.1 qb:AF065485.1 db:NM 003794 1 |
| 218194_at | small fragment all fragment nu |
| 220560_at | SMS3 protein (SMS3), mRNA. /FEA-mRNA /GE |
| 208270_s_at | gb:NM 020216.2 /DEF=Homo sapiens arginyl aminopeptidase (aminopeptidase B) (RNPEP), mRNR. /FEA-mRNR /GEN-RNPEP /PROD-arginyl aminopepti-, dase (aminopeptidase B) /DB XREF-gi:13443030 /UG=Hs.283667 arginyl aminopeptidase (aminopeptidase R) /Fi=ch-NR 070316 2 |
| 220175_s_at |), mRNA. /FEA=mRNA |
| 212429_s_at | NE=IMAGE: 25775 |

| 209058_at | ns mRNA for hMBFlalpha, complete cds. /FEA=mRNA /PROD=h r 1 /FL=gb:NM 003792.1 gb:AB002282.1 |
|-------------|--|
| 212535_at | Consensus includes gb:AA142929 /FEA-EST /DB_XREF-gi:1712307 /DB_XREF-est:z140g07.s1 /CLONE-IMAGE:504444 /UG=Hs.288993 ESTs |
| 201524_x_at | ens ubiquitin-conjugating enzyme EZN (homologous to yeast UBC13) (UBEZN), mRNA. /FEA-mRNA /GEN- zyme EZN (homologous toyeast UBC13) /DB_XREF=g1:4507792 /UG-Hs.75355 ubiquitin-conjugating enzym 83004.1 gb:BC000396.1 gb:BC003365.1 gb:RN 003348.1 |
| 201716_at | ens sorting nexin 1 (SNX1), mRNA. /FEA-mRNA /GEN-SNX1 /PROD-sorting nexin 1 /DB_XREF-g1:4507138 57.1 qb:U53225.1 qb:AF065483.1 qb:NN 003099.1 |
| 213097_s_at | 7 /FEA-EST /DB_XREF-g1:4075764 /DB_XREF-est:qq28f09.xl /CLONE-IMAGE:1933865 /UG-Hs.82254 zuotin relate |
| 213532_at | 1 /CLONE=IMAGE:2386986 /UG-H |
| 210766_s_at | ns trachea cellular apoptosis susceptibility protein (CSE1) mRNA, cc eptibility protein /DB_XREF-gi:3560554 /UG-Hs.90073 chromosome segre |
| 209523_at | l /DEF-Homo sapiens cDNA FLJ10756 fis, A /DB XREF-gi:7022983 /UG-Hs:122752 TA 0701.1 gb:AE057694.1 gb:NM 003184.1 |
| 209276_s_at | gb:AF162769.1 /DEF-Homo sapiens thioltransferase mRNA, complete cds. /FEA-mRNA /PROD-thioltransferase /DB_XREF-g1:5442445 /UG-Hs.28988 glutaredoxin (thioltransferase) /FL-gb:BC005304.1 gb:AF162769.1 gb:D21238.1 |
| 200848 at | 38 /E |
| 212406_s_at | 13.1 /DEF-Homo sapiens mRNA for KIAA1050 protein, partial cds: /FEA-mRNA / 1628 hypothetical protein FLJ10883 |
| 212072_s_at | 51 /DEF=Human DNA sequence from clone RP5-863C7 on chromosome 20p12 (EC 2.7.1.37), ESTs, STSs and GSSs /FEA-mRNA_2 /DB_XREF=gi:5738437 |
| 203159_at | 5.1 /DEF-Momo sapiens :AF327434.1 gb:AB02064 |
| 203521_s_at | ens endocrine regulator (HRIHFB2436), mRNA. /FEA-mRNA /GEN-HRIHFB2436 /PROD-endocrine rec 1433 endocrine regulator /FL-gb:AF121141.1 gb:NM 014345.1 |
| 204439_at | ens hypothetical protein, expressed in osteoblast (G33686), mRNA. /FEA=mRNA /GEN=GS3686 /PROD- ast /DB_XREF=g1:5803026 /UG=Hs.75470 hypothetical protein, expressed in osteoblast /FL=gb:AB000 |
| 218646_at | lens hypothetical protein ELJ20534 (FLJ20534), mRNA. /FEA-mRNA /GEN-FLJ20534 /PROD-hypotheti /UG-Hs.44344 hypothetical protein FLJ20534 /FL-gb:AL136673.1 gb:NA 017867.1 |
| 213168_at | FEA-EST /DB_XREF-g1:11006526 /DB_XREF-est:AU145005 /CLONE-HEMBA1003603 /UG-Hs.44450 |
| 210371_s_at | ns, Similar to retinoblastoma binding protein 4, clone MGC:1399 btein 4 /DB XREE—gi:131111850 /UG-Hs.16003 retinoblastoma-binding |
| 200052_s_at | enhancer binding factor 2, 45kD (ILF2), mRNA. / UG-Hs.75117 interleukin enhancer binding factor 2 |
| 201019_s_at | gb:NM 001412.1 /DEF-Homo sapiens eukaryotic translation initiation factor lA (EIFIA), mRNA. /FEA=mRNA /GEN-EIFIA /PROD=eukaryotic transla- tion initiation factor lA /DB_XREF-gi:4503498 /UG-Hs.4310 eukaryotic translation initiation factor lA /FL-gb:BC000793.1 gb:Ll8960.1 gb:NM 001412.1 |
| 219892_at | ens transmembrane 6 superfamily member 1 (TW6SF1), mRNA. /FEA-mRN 3194198 /UG-Hs.133865 transmembrane 6 superfamily member 1 /FI-gb |
| 218095_s_at | <pre>gb:NM 018475.1 /DEF=Homo sapiens uncharacterized hypothalamus protein HTMP (LOC3585b), mRNA. /FEA=mRNA /GEN=LOC3585b /FRUD=uncharacterized hypothalamus protein HTMP /DB_XREF=gi:8923860 /UG=Hs.236510 uncharacterized hypothalamus protein HTMP /EL=gb:BC003545.1 gb:AF220188.1 gb:NN 018475.1 gb:AF183409.1</pre> |

| <u> </u> | at | gb:NM 016129.1 /DEF=Homo sapiens COP9 complex subunit 4 (LOC51138), mRNA. /FEA=mRNA /GEN=LOC51138 /PROD=COP9 complex subunit 4 (LOC51138), mRNA. /FEA=mRNA /GEN=LOC51138 /PROD=COP9 complex subunit 4 /FL=gb:BCO04302.1 gb:AF100757.1 gb:NM 016129.1 |
|------------|-------------|--|
| <u>(1)</u> | , | .6 /FEA=EST /DB_XREF=gi:12428927 /DB_XREF=est 146.1 |
| [2] | | |
| 10 | | 99 /FEA-EST /DB_XREF-g1:42 |
| ~_ | | Consensus includes gb:AW161626 /FEA=EST /DB XREF=g1:6300659 /DB XREF=est:au68bll.xl /CLONE=IMAGE:2781405 /UG-Hs.21739 Homo sapiens mRNA, CDN DKF2p58611518 (from clone DKF2p58611518) |
| (0) | _at . | gb:BE349584 |
| 101 | 208174_x_at | gb:NM 005089.1 /DEF-Homo sapiens U2 small nuclear ribonucleoprotein auxiliary factor, small subunit 2 (U2AFIRS2), mRNA. /FEA-mRNA /GEN-U2AFIRS2 / |
| | | PROD=U2 small nuclear ribonucleoprotein auxiliaryfactor, small subunit-related protein 2 /DB_XREF=gi:4827045 /UG=Hs.171909 U2 small nuclear ribonucleoprotein auxiliary factor, small subunit 2 /FL=qb:DH9677.1 qb:NM 005089.1 |
| īΩ. | 53912_at | Cluster Incl. AI668643:zb13f10.x5 Homo sapiens cDNA, 3 end /clone=IMAGE-301963 /clone_end=3 /gb=A1668643 /gi=4827951 /ug=Hs.15827 /len=601 |
| 2_ | | Consensus includes gb:AA156777 /FEA-EST /DB_XREF-g1:1728392 /DB_XREF-est:z118c08.s1 /CLONE-IMAGE:502286 /UG-Hs.25431 KIAA1219 protein |
| | | ns deubiquit 68 ubiquitir |
| 7 | a t | <pre>general transcription factor I ription factor IIIC, polypeptid (beta subunit, 110kD) /FL=db:DI</pre> |
| 10 | 212584_at | 9 /FEA=EST /DB_XREF=gi:12 |
| <u> </u> | | Consensus includes gb:AB014548.1 /DEF=Homo sapiens mRNA for KIAA0648 protein, partial cds. /FEA=mRNA /GEN=KIAA0648 /PROD=KIAA0648 protein / DB XREE=gi:3327109 /UG=Hs.31921 KIAA0648 protein |
| | | gb:NM 018130.1 /DEE-Homo sapiens hypothetical protein FLJ10539 (FLJ10539), mRNA. /FEM-mRNA /GEN-FLJ10539 /PROD-hypothetical protein FLJ10539 /DB XREF-gi:8922499 /UG-Hs.93391 hypothetical protein FLJ10539 /FL-qb:NM 018130.1 |
| ici T | | 26.1 /DEF=Hom othetical pro |
| (7) | 216194_s_at | ng CAPNS and POLZRI, genom |
| 10 | _s_at | 20 en |
| <u> </u> | | 38 /FEA-EST /DB_XREF-g1:11947783 /DB_XREF-esc |
| <u>N</u> | | 9b:NM 016562.1 /DEF=Homo sapiens toll-like receptor 7 (LOC51284), mRNA. /FFA=mRNA /GEN=LOC51284 /PROD=toll-like receptor 7 /FL=qb:AF240467.1 qb:NN 016562.1 qb:AF245702.1 |
| (1) | | gb:NM 019018-1 /DEF-Homo sapiens hypothetical protein (FLJ11127), mRNA. /FER-mRNA /GEN-FLJ11127 /PROD-hypothetical protein /DB XREF-gi:9506640 /UG-Hs:91165 hypothetical protein /FL-qb:NM 019018.1 |
| | 213229_at | Consensus includes gb:BF590131 /FEA-EST /DB_XREF-gi:11682455 /DB_XREF-est:nab19e04.x1 /CLONE-IMAGE:3266383 /UG-Hs.87889 helicase-moi |
| [0] | at | Consensus includes gb:AI272899 / FEA-EST / DB_XREF=gi:3895167 / DB_XREF-est:q147c07.x1 / CLONE-IMAGE:1875468 / UG-Hs.3852 KIAA0368 protein |
| | | Consensus includes 9b:AB028999.1 /DEF=Homo sapiens mRNA for KIAA1076 protein, partial cds. /FEA=mRNA /GEN=KIAA1076 /PROD=KIAA1076 protein / DB XREF=gi:5689488 /UG-Hs.154525 KIAA1076 protein |
| <u>~ 1</u> | at | DC41 m gb:NM |
| <u>~</u> | 221522_at | gb:AL136784.1 /DEF-Homo sapiens mRNA; cDNA DKF2p434L0718 (from clone DKF2p434L0718); complete cds. /FEA=mRNA /GEN-DKF2p434L0718 //PROD-hypothetical |

| • | |
|-------------|---|
| | protein /DB_XREF=gi:12053080 /UG-Hs.59236 Homo sapiens mRNA; cDNA DKFZp434L0718 (from clone DKFZp434L0718); complete cds /FL-gb:AL136784.1 |
| 205930_at | gb:NN 005513.1 /DEF-Homo sapiens general transcription factor IIE, polypeptide 1 (alpha subunit, 56kD) (GTF2E1), mRNA. /FFA-mRNA |
| | PROD-general transcription factor IIE, polypeptide 1(alpha subunit, 56kD) /DB_XREF-gi:5031726 /UG-Hs.145381 general transcription factor |
| | |
| 210312_s_at | F=Homo sapie 12803610 /UC |
| 213838_at | 6 /FEA=EST /DB_XREF=gi:1780105 /DB_XREF=est:zp83g09.s1 /CLONE=IMAGE:626848 /UG=H |
| 202520_s_at | ens mutī (E. coli) homolog l (colon cancer, nonpolypos ,/UG-Hs.57301 mutī (E. coli) homolog l (colon cancer, |
| 221834_at | ST /DB_XREF-g1:10302103 /DB_XREF-est:AV700132 /CLONE-GKCGSE03 /UG-Hs.295923 seven in absentia |
| 210942_s_at | ıns mRNA for alpha2,3-sialyltransferase ST3Gal VI, complete cds. /FEA-mRNA /GEN-ST3Gal VI /PRC 'DB XREF-qi:4827246 /UG-Hs.34578 alpha2,3-sialyltransferase /FL-gb:AB022918.1 |
| 202396_at | ens transcription factor CA150 (CA150), mRNA. /FEA÷mRNA /GEN÷CA150 /FF 1063 transcription factor CA150 /FL=9b:NF017189.1 gb:NM 006706.1 |
| 219598_s_at | -mRNA /GEN=PTD013 |
| 202502_at | 7 |
| 211968_s_at | 3 /FEA-EST /DB_XREF-gi:5755646 /DB_XREF-est:wt25a07.x1 /CLONE-INAGE:250846 5348.1 |
| 218027_at | ens HSPC145 protein (HSPC145), mRNA. /FEA-mRNA /GEN-HSPC145 /PROD-HSPC145 protein /DB_XREF-gi:76 /FI-gb:AL136665.1 gb:BC000891.1 gb:AF161494.1 gb:NM 014175.1 |
| 217792_at | mRNA /GEN=SNXS /PROD=sor |
| 221036_s_at | gb:NM 031301.1 /DEF-Homo sapiens hypothetical protein DKFZp564D0372 (DKFZp564D0372), mRNA. /FEA-mRNA /GEN-DKFZP564D0372 / |
| 201723_s_at | |
| 202351_at | 3432555 /D tigen CD51 |
| 208853_s_at | gb:L18887.1 /DEF=Human calnexin mRNA, complete cds. /FEA=mRNA /PROD=calnexin /DB_XREF=gi:306480 /UG=Hs.155560 calnexin /FL=gb:NN_001746.1 qb:BC003552.1 qb:M98459.1 qb:M98452.1 gb:L10284.1 gb:L18887.1 |
| 213278_at | Consensus includes gb:RW014788 /EEA-EST /DB_XREF-gi:5863545 /DB_XREF-est:UI-H-BIO-aae-h-10-0-UI.s1 /CLONE=IMAGE:2709354 /UG-Hs.48802 Homo |
| 203447_at | |
| 212459_x_at | 40 /FEA-EST /DB_XREF-q1:11686264 /DB_XREF-est:nab48f10.x1 /CLON bunit |
| 218514_at | hypothetical protein FLJ10587 (FLJ10587), mRNR. /FEA-mRNR /GEN-FLJ1058 Hs.7296 hypothetical protein FLJ10587 /FL-gb:NN 018149.1 |
| 212648_at | viens mRNA full length insert cDNA clone EUROINAGE 48814. /FEA=mRNA /PROD=hypothetica) ise A Arabidopsis thaliana /DB XREF=gi:5102732 /UG=Hs.95665 hypothetical protein |
| 200708_at | <pre>gb:NM 002080.1 /DEF=Homo sapiens glutamic-oxaloacetic transaminase 2, mitochondrial (aspartate aminotransferase 2) (GOT2), nuclear gene encoding mitochondrial protein, mRNA. /FEA=mRNA /GEN=GOT2 /PROD=aspartate aminotransferase 2 precursor /DB_XREF=g1:4504068 /UG=Hs.170197, glutamic-oxaloacetic transaminase 2, mitochondrial (aspartate aminotransferase 2) /FI=gb:BC000525.1 gb:M2c632.1 gb:NM 002080.1</pre> |

| 201064_s_at | tens poly(A)-binding protein, cyroplasmic 4 (inducible form) (PABPC4), mRNA. / 1, cytoplasmic 4(inducible form) /DB_XREF=gi:6552335 /UG=Hs.169900 poly(A)-bind 19.2 |
|---------------|---|
| 218604_at | ens integral inner nuclear membrane protein (MANI), mRNA. /FEA-mRNA /GEN-MAN 7706606 /UG-Hs.7256 integral inner nuclear membrane protein /FL-gb:AFI12299. |
| 214864_s_at . | Consensus includes gb:AK024386.1 /DEF-Homo sapiens cDNA FLJ14324 fis, clone PLACE4000100, highly similar to Homo sapiens hydroxypyruvate reductase(GRHPR) gene. /FEA-mRNA /DB XREF-gi:10436760 /UG-Hs.155742 glyoxylate reductasehydroxypyruvate reductase |
| 202717_s_at | ens C erevi |
| 201643_x_at | 9b:NH 016604.1 /DEF=Homo sapiens putative zinc finger protein (LOC51780), mRNA. /FEA-mRNA /GEN-LOC51780 /PROD-putative zinc finger protein /DB XREF=gi:7706598 /UG-Hs.24125 putative zinc finger protein /FI-gb:AF251039.1 gb:NN 016604.1 |
| 213803_at | Consensus includes gb:BG545463 /FEA=EST /DE_XREF=gi:13544128 /DB_XREF=est:602572695F1 /CLONE-IMAGE:4701118 /UG=Hs.180446 karyopherin (im- portin) beta l |
| 204689_at | gb:NN 001529.1 /DEF=Homo sapiens hematopoietically expressed homeobox (ННЕХ), mRNA. /FEA-mRNA /GEN=HHEX /PROD-hematopoietically expressed homeobox /DB XREF=g1:10835016 /UG=Hs.118651 hematopoietically expressed homeobox /FE=gb:NM 001529.1 gb:L16499.1 gb:NN 002729.1 |
| 218228_s_at | ens tankyrase 2 (TNKL), mRNA. /FEA-mRN F329696.1 gb:NM 025235.1 gb:AF342982.1 |
| 202918_s_at | sapiens CGI-95 protein mRNA, complete cds. /FEA-mRNA /PROD=CGI-95 protein /DB_XREF-g1:4929658 /UG-Hs.1 -gb:AB015441.1 gb:BC005237.1 gb:AF151853.1 gb:AL080070.1 gb:NH 015387.1 |
| 212263_at | Consensus includes gb:AII14716 /FEA=EST /DB XREF=gi:6360061 /DB_XREF=est:HA1315 /UG-Hs.15020 homolog of mouse quaking QKI (KH domain RNA binding protein)/FL=gb:AF142419.1 gb:AF142472.1 |
| 221825 at | 93 /FEA-EST /DB_XREF-gi:11013314 /DB_XREF-est:AU151793 /CLONE-NT2RP2006115 /UG-Hs.157078 H 2033 |
| 211047_x_at | gb:BC006337.1 /DEF-Homo sapiens, clone MGC:12798, mRNA, complete cds. /FEA-mRNA /PROD-Unknown (protein for MGC:12798) /DB_XREF-g1:13623468 /FL-qb:BC006337.1 |
| 209798_at | IRNA, complete cds. /FEA-mRNA /GEN-NPAT /DB_XREF-gi:1304113 /UG-Hs.89385 nuclear protein, ataxia-)02519.1 |
| 204361_s_at | 9 - |
| 218127_at | Consensus includes gb:AI804118 /FEA=EST /DB XREF=gi:5369590 /DB XREF=est:tc68e01.x1 /CLONE=IMAGE:2069784 /UG-Hs.84928 nuclear transcription factor Y, beta /FL-gb:NM 006166.2 gb:BC005316.1 gb:BC005317.1 gb:L06145.1 |
| 220355_s_at | hypothetical protein FLJ10645 (FLJ10645), mRNA. HS 44143 polybromo 1 /FL=gb:AF177387:1 gb:NM 018 |
| 202930_s_at | gb:NM 003850.1 /DEF=Homo sapiens succinate-CoA ligase, ADP-forming, beta subunit (SUCLA2), mRNA. /FEA=mRNA /GEN=SUCLA2 /PROD=succinate-CoA ligase, ADP-forming, beta subunit /FL=gb:NN_003850.1 gb:AB035863.1 |
| 203745_at | Consensus includes gb:A1801013 /FEA=EST /DB XREF=gi:5366485 /DB XREF=est:wg15d09.x1 /CLONE=INAGE:2365169 /UG=Hs.211571 holocytochrome c Synthase (cytochrome c heme-lyase) /FL-gb:U36787.1 gb:NM 005333.1 |
| 200597_at | :0 /FEA=EST /DB_XREF=gi:3539086 /DB_XREF=est:qa48al0. punit 10 (theta, 150170kD) /FL=gb:D50929.1 gb:U58046. |
| 218519_at | gb:NM 017945.1 /DEF=Homo sapiens hypothetical protein FLJ20730 (FLJ20730), mRNA. /FEA=mRNA /GEN=FLJ20730 /PROD=hypothetical protein FLJ20730 /DB XREF=gi:8923656 /UG-Hs.237480 hypothetical protein FLJ20730 /FL=gb:BC005207.1 gb:NM 017945.1 |
| 203947_at | gb:NN 001326.1 /DEF=Homo sapiens cleavage stimulation factor, 3 pre-RNA, subunit 3, 77kD (CSTF3), mRNA. /FEA-mRNA /GEN-CSTF3 /PROD=cleavage stimulation factor subunit 3 /DB_XREF=gi:4557494 /UG-Hs.180034 cleavage stimulation factor, 3 pre-RNA, subunit 3, 77kD /FL-gb:NN 001326.1 gb:U15782.1 |
| 219002_at | |
| 220122_at | nRNA. /FEA-mRNA /GEN-FLJ22344 /PROC FL-gb:NY 024717.1 |
| 203882_at | gb:NM 006084.1 /DEF=Homo sapiens interferon-stimulated transcription factor 3, gamma (48kD) (ISGF3G), mRNA. /FEA=mRNA /GEN=ISGF3G / PROD=interferon-stimulated transcription factor 3,gamma (48kD) /DB XREF=gi:5174474 /UG=Hs.1706 interferon-stimulated transcription factor |

| | 3, gamma (48kD) /FLI-gb:M87503.1 gb:NM_006084.1 |
|-------------|---|
| 212174_at | Consensus includes gb:W02312 /FEA-EST /DB_XREF-gi:1274291 /DB_XREF-est:za08d08.rl /CLONE-IWAGE:291951 /UG-Hs.171811 adenylate kinase 2 |
| 202567_at | н – н |
| 218557_at | 1910459 /UG=Hs |
| 211976_at | Consensus includes gb:AK026168.1 /DEF-Homo sapiens cDNA: FLJ22515 fis, clone HRC12122, highly similar to AF052101 Homo sapiens clone 23872 mRNA sequence. /FEA-mRNA /DB XREF-g1:10438931 /UG-Hs.188882 Homo sapiens cDNA: FLJ21862 fis, clone HEP02321, highly similar to AF052101 Homo sapiens clone 23872 mRNA sequence |
| 201990_s_at | <pre>db:NM 001310.1 /DEF=Homo sapiens cAMP responsive element binding protein-like 2 (CREBL2), mRNM. /FEA=mRNM /GEN=CREBL2 /PROD=CAMP respon- sive element binding protein-like 2 /DB_XREF=g1:4503034 /UG=Hs.13313 CAMP responsive element binding protein-like 2 /FL=gb:AF039081.1 qb:NM 001310.1</pre> |
| 208956 x at | EF-Human deoxyuridine triphosphatase (DUT) mRNA, complete cds. /FEA-mRNA /GEN-DUT /PROD-deoxyuridine triphosph 21817 /UG-HS.82113 dUTP.pyrophosphatase /FL-gb:AB049113.1 gb:U31930.1 gb:UG2891.1 gb:M89913.1 gb:NM 001948.1 |
| 213939_s_at | 41 /FEA=EST /DB_XREF=g1:5545690 /DB_XREF=est:we28f04.xl /CLONE=INAGE:2342431 /UG-Hs.306623 Homo sapi 1668 |
| 201198_s_at | 31 /FEA-EST /DB_XREF-gi:5514047 /DB_XREF-est:w113h07.x1 /CLONE-IMAGE:242. ATPase, 1 /FL-gb:D44466.1 gb:Nh 002@07.1 |
| 203010_at | gb:NM 003152.1 /DEE-Homo sapiens signal transducer and activator of transcription 5A (STAT5A), mRNA. /FEA-mRNA /GEN-STAT5A /PROD-signal transducer and activator of transcription 5A /DB_XREF-gi:4507256 /UG-Hs.167503 signal transducer and activator of transcription 5A /FL-ab:U43185.1 gb:NM 003152.1 gb:L41142.1 |
| 218067_s_at | |
| 219077_s_at | gb:NH 016373.1 /DEE-Homo sapiens WM domain-containing oxidoreductase (VMOX), mRNA. /FEA-mRNA /GEN-WVOX /PROD-FOR II /DB_XREF-gi:7706522 / UG-Hs.519 WW domain-containing oxidoreductase /FL-gb:AF211943.1 gb:NM 016373.1 gb:AF227527:1 |
| 215424_s_at | DB_XREF=est:AV689564 /CLONE=GKCEAAO6 /UG=Hs.79008 SKI-INTER |
| 219043_s_at | gb:NN 024065.1 /DEE-Homo sapiens hypothetical protein MGC3062 (MGC3062), mRNA. /FEA-mRNA /GEN-MGC3062 /PROD-hypothetical protein MGC3062 / DB XREF-gi:13129043 /UG-Hs.94576 hypothetical protein MGC3062 /FEL-gb:AF267853.1 gb:BC001021.1 gb:NM 024065.1 |
| 209313_at | gb.AB044661.1 /DEF-Homo sapiens XAB1 mRNA for XPA binding protein 1, complete cds. /FEA-mRNA /GEN-XAB1 /PROD-XPA binding protein 1 / DB XREF-gi:11094140 /UG-Hs.18259 XPA binding protein 1; putative ATP(GTP)-binding protein /FL-gb.AB044661.1. |
| 202944_at | |
| 208985_s_at | lens, eukaryotic t translation initie bunit 1 (alpha, 3 |
| 211656_x_at | |
| 209358_at | ജ I |
| 207431_s_at | iens degenerative spermatocyte (homolog Drosophila; lipid desaturase) (DEGS), mRNA. /FEA=mRNA /GEN=DE yte /DB_XREF=gi:4505192 /UG=Hs.185973 degenerative spermatocyte (homolog Drosophila; lipid desaturase) 76.1 |
| 213070_at | 36 /FEA-EST /DB_XREF=g1:10284299'/DB_XREF=est:AV682436 /CLONE-GKBABE08 /UG-Hs "2p564L222) |
| 201111_at | <pre>lens brain cellular apoptosis suscept sceptibility protein /DB_XREF=q1:3560 .1 qb:NN 001316.1</pre> |
| 204370_at | gb:NN 006831.1 /DEF-Homo sapiens ATPGTP-binding protein (HEAB), mRNA. /FEA-mRNA /GEN-HEAB /PROD-ATPGTP-binding protein /DB_XREF-g1:5803028 /UG-HS.87465 ATPGTP-binding protein /FL-gb:BC000446.1 gb:U73524.1 gb:NN 006831.1 |
| | |

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|-------------|---|
| 202911_at | gb:NM 000179.1 /DEF-Homo sapiens muts (E. coli) homolog 6 (MSH6), mRNA. /FEA-mRNA /GEN-MSH6 /PROD-muts (E. coli) homolog 6 /FL-gb:U28946.1 gb:BC004246.1 gb:NM 000179.1 gb:U54777.2 |
| 204197_s_at | gb:NM 004350.1 /DEE-Homo sapiens runt-related transcription factor 3 (RUNX3), mRNA. /FEA-mRNA /GEN-RUNX3 /PROD-runt-related transcription factor 3 /FL-gb:NM 004350.1 |
| 217809_at | lens HSPC028 protein (HSPC028), mRNA, /FEA-mRNA /GEN-HSPC028 /PROD-HSPC028 protein /DB_XREF- /FL-gb:AF110323.1 |
| 201054 at | 9 /FER-EST /DB XREF-g1:11772191 /DB XREF-est:601660283R1 /CLONE-IMAGE:3906044 /UG-Hs.77492 hei L-gb:BC001008.1 gb:NN 006805.1 gb:UZ3803.1 |
| 200994_at | 7 /FEA-EST /DB_XREF-gi:13050002 /DB_XREF-est:602386007F1 /CLONE-IMAGE:4515 391.1 |
| 201572_x_at | ens dCMP deaminase (DCTD), mRNA. /FEA=mRNA /GEN=DCTD 1 gb:NM 001921.1 |
| 218195_at | ens hypothetical protein FLJ12910 (FLJ12910) |
| 219905_at | DB XREF-gi:13375745 /UG-Hs.15929 hypothetical procein FL012910 /FL-go:NW U245/5.1 gb:NM O18538 |
| 209362_at | 0 /FEA-EST /DB XREF-q1:4899874 /DB XREF-est:wd39h05.x1 /C loq /FL-qb:U46837.1 qb:U52960.1 qb:NM 004264.1 |
| 203983_at | translin-associated factor > |
| 212074_at | 4 /FEA=EST /DB_XREF=gi:10586110 / |
| 204859_s_at | gb:NM_013229.1 /DEF-Homo sapiens apoptotic protease activating factor (APAFI), transcript variant 1, mRNA. /FEA-mRNA /GEN-APAFI /PROD-apoptotic protease activating factor isoform a /DB_XREF-gi:7108332 /UG-Hs.77579 apoptotic protease activating factor /FL-qb:AB007873.1 gb:AF134397.1 gb:NM 013229.1 |
| 205707_at | ens interleukin 17 receptor (IL17R), mRNA. /FEA=mRNA /GEN=IL17R /PROD- 9751 interleukin 17 receptor /FL-gb:U58917.1 gb:NN 014339.1 |
| 218716_x_at | iens CGI-02 protein (CGI-02), mRNA./FEA-mENA /GEN-CGI-02 /PROD-CGI-02 protei 22.1 gb:AF132937.1 gb:NM 012123.1 |
| 217940_s_at | ens hypothetical protein FLJ10769 (FLJ10769), mRNA. /FEA-mRNA /UG-Hs:8083 hypothetical protein FLJ10769 /FL-gb:AF151071.1 gb |
| 219802_at | ens hypothetical protein FLJ22028 (FLJ22028), mRNA. /FEA-mRNA /GEN-FLJ22028 /PROD-hypo //UG-Hs.192570 hypothetical protein FLJ22028 /FL-gb:NM 024854.1 |
| 209259_s_at | REF |
| 212037_at | Consensus includes gb:BF508848 /FEA-EST /DB_XREF-gi:11592146 /DB_XREF-est:UI-H-BI4-aor-e-06-0-UI.s1 /CLONE-IMAGE:3085907 /UG-Hs.44499 Dinin, desmosome associated protein |
| 211063_s_at | protein |
| 203791_at | x-like 1 (DKKL1), mRNA. /FEA-mRNA /GEN-DMKL1 /PROD-Dmx-like 1 /DB_XREF-q::9961348 |
| 203405_at | ens Down syndrome critical region gene 2 (DSCR2), mRNA. /FEA-mRNA / 4505022 /UG-Hs.5198 Down syndrome critical region gene 2 /FL-qb:BCO |
| 201604_s_at | lens myosin phosphatase, target subunit 1 (MYFT1), mRNA. /FEA-mRNA /GEM-MYPT1 /PROE 5 /UG-Hs.16533 myosin phosphatase, target subunit 1 /FL-gb:NM 002480.1 |
| 218348_s_at | ens HSPCO55 protein (HSPC055), mRNA. /FEA-mRNA /GEN-HSPC055 /PROD-HSPC055 protein /DB_XREF-gi:7661761 , /FL-gb:AF161540,1 gb:NM 014153.1 |
| 205659_at | iens histone deacetylase 7B (HDAC7B-PENDING), mRNA. /FEA-mRNA /GEN-HDAC7B-PENDING /PROD 16753 histone deacetylase 7B /FI-gb:AB018287.1 gb:NN 014707.1 |
| 202268_s_at | gb:NM 003905.1 /DEF=Homo sapiens amyloid beta precursor protein-binding protein 1, 59kD (APBBPI), mRNA. /FEA=mRNA /GEN=APPBPI /PROD=Amyloid beta precursor protein-binding protein! /DB XREF=gi:4502168 /UG-Hs.61828 amyloid beta precursor protein-binding protein 1, |

| 202399_s_at | adaptor-related protein complex 3, sigma 2 subunit (AP3S2), mRNA. /FEA-mRNA /GEN- 2subunit /DB_XREF-g1:5031580 /UG-Hs.154782 adaptor-related protein complex 3, sigm |
|-------------|--|
| 218515_at | ans hypothetical protein (LOC51325), mRNA. /FEA=mRNA /GEN=LOC51325 /PROD=hypothetical protein 161 hypothetical protein /FL=gb:AF2208862.1 gb:NM 016631.1 |
| 209180_at | geranyl transferase type II beta-subunit mRMA, complete cds. /FEA-mRMA./PRUD-geranylgeranyl transf 16503 /UG-Hs.78948 Rab geranylgeranyltransferase, beta subunit /FL-biU49245.1 gb:NM 004582.1 |
| 205105_at | mannosidase, alpha, class:2A, member 1 (MANZA1), mRNA. /FEA-mR 158697 /UG-Hs.32965 mannosidase, alpha, class 2A, member 1 /FL- |
| 200666_s_at | s heat shock 5 DnaJ (Hsp4 |
| 200972_at | NA, complete cds. /FEA-mRNA./PROD-tetraspan 3 / UB_XKEF-g1:126 054840.1 gb:NM 005724.1 gb:AF133423.1 |
| 218352_at | =mRNA /GEN=ELJ10/16 /PROD=hypothetical pro 191.1 |
| 217728_at | ens S100 calcium-binding protein A6 (calcyclin) (S100A6), mRNA. /FEA=mRNA /GEN=5100A6 /EROD=5100 calci 1:9845517 /UG=Hs.275243 S100 calcium-binding protein A6 (calcyclin) ·FE=Gp:BC001431.1 gb:NM 014624.2 |
| 211971_s_at | 3 /FEA=EST /DB_XREF=gi::4737587:/I |
| 212500_at | 9.1 /DEF=Homo sapiens mRNA; cDNA DKFZp564C046 (from clone DKF2p564C046). /FEA=mRNA /DB_XREF=91: A; cDNA DKF2p564C046 (from clone DKF2p564C046) |
| 218473_s_at | hypothetical protein FLJ22329 (FLJ22329), mRNA. /FFA=mRNA /GEN=FLJ22329 /PROD=hypothetical prote: 5-Hs.61478 hypothetical protein FLJ22329 /FL=qb:NM 024656.1 |
| 203580_s_at | db:NM 003983.1 /DEE=Homo sapiens solute carrier family 7 (cationic amino acid transporter, y+ system), member 6 (SLC7A6), mRNA. /FEA-mRNA /GEN-SIC7A6 /PROD-solute carrier family 7 (cationic amino acidtransporter, y+ system), member 6 /DB XREF-gi:4507052 /UG-Hs.10315 solute |
| 200900_s_at | Consensus includes gb:AI583537 /FEA-EST /DB XREF-gi:4569434 /DB XREF-est:ts12d03.x1 /CLONE-IMAGE:2228357 /UG-Hs.75709 mannose-6-phosphate receptor (cation dependent) /FL-gb:NN 002355.2 gb:NH 5985.1 |
| 221652_s_at | ns PNAS-25 mRNA, complete cds. /FEA=mRNA /PROD=FNAS-25 /DB_XREF=gi:12751064 /UG=Hs.22595 hypoth |
| 217750_s_at | apiens hypothetical protein FLJ13855 (FLJ13855), mRNA. /FEA=mRNA /GEN=FLJ13855 /PROD=hypothetical p .168232 hypothetical protein FLJ13855 /FL-qb:NM 023079.1 |
| 203544_s_at | signal transducing adaptor molecule (SH3 domain and ITAM motif) 1 (ST molecule (SH3 domainand ITAM motif) 1 /DB_XREF=gi:4507248 /UG=Hs.1534 1 /FL=gb:U43899.1 gb:NM 003473.1 |
| 221580_s_at | ns, clone MGC:5306, mRNA, complete cds. /FEA-mRNA /PROD-Unknown (protein for MGC:5306) /DB_XREF- otein MGC5306 /FL-gb:BC001972.1 |
| 202629_at | Consensus includes gb:AV681579 /FEA=EST /DB XREF=g1:10283442 /DB XREF=est:AV681579 /CLONE=GKBAFEUS /UG=HS:84044 amylold Deta precursor protein (cytoplasmic tail)-binding protein 2 /FL=gb:AF017782.1.gb:NM 006380.1 |
| 205763_s_at | mkNA /GEN=DUXIS x polypeptide l |
| 212337_at | 8 /FEA=EST /DE |
| 212244_at | Consensus includes gb:AL050091.1 / DEF-Homo saptens mRNA; cDNA DRE2p38611918 (Irom clone DRE2p3861918); partial cds. / FEA-WHKWA / GEN-DRE2p58611918 / PROD-hypothetical protein / DB XREF-941:4084111 / VG-Hs 5283 DRE2F3861918 protein / DB XREF-941:4084111 / VG-Hs 5283 DRE2P3861918 protein / DB XREF-941:4084111 / VG-Hs 7-203-66 11 / OF/ANT-194036 / VG-Ms |
| 222011_s_at | ST /DB XREF=g1:11131299 /DB XREF=est://d83803.X1 / CLONE=IMAGE:3/04930 /OG=ns.z/10944 dretyl-Coenzyme zyme A thiolase) |
| 204725_s_at | gb:NM 006153.1 /DEF=Homo sapiens NCK adaptor protein 1 (NCK1), mkNA. /FEA=mKNA / GEN=NCK1 / FKUD=NCA adaptor protein 1 /DD_AREF=91:3433733 / UG-Hs.54589 NCK adaptor protein 1 /FL=qb:NM 006153.1 |

| 208860_s_ar | gb:U09820.1 /DEE-Human helicase II (RAD54L) mRNA, complete cds. /FEA-mRNA /GEN-RAD54L /PROD-helicase II /DB XREF-g1:606832 /UG-Hs.96264 31pha thalassemiamental retardation syndrome X-linked (RAD54 (S. cerevisiae) homolog) /FL-db:U09820.1 ch:NN-000489 1 ch:U72037 2 |
|-------------|---|
| 203224_at | 12037283F1 /CLONE=IMAGE:4185212 /UG-Hs.375 |
| 208619_at | gb:L40326.1 /DEF=Homo sapiens Hepatitis B virus X-associated protein 1 mRNA, complete cds. /FEA=mRNA /PROD=X-associated protein 1 |
| | UG-Hs.108327 damage-specific DNA binding protein 1 (127kb) /FL-gb:U18299.1 gb:U32986.1 gb:NM 001923.2 gb:L40326.1 |
| 208765_s_at | Consensus includes gb:NM 005826.1 /DEF=Homo sapiens heterogeneous nuclear ribonucleoprotein R (HNRRR), MRNA, /FEA-CDS /GEN-HNRPR PROD-Intercognenous nuclear ribonucleoprotein R /DB_XREF=gi:5031754 /UG-Hs.15265 heterogeneous nuclear ribonucleoprotein R |
| 220832_at | : G |
| | UG-HS.272410 Toll-like receptor 8 /FL-qb:AF246971.1 db:NM 016610.1 db:AF245703.1 |
| 208398_s_at | |
| 205077_s_at | db:NM_002643.1 /DEF-Homo sapiens phosphatidylinositol glycan, class F (PIGE), mRNA. /FER-mRNA /GEN-PIGE /PROD-phosphatidylinositol glycan, class F /FI-ch:D13435 1 ob:NM_00543 1 |
| 210616_s_at | lens mRNA for KIAA0905 protein, complete cds. /FEF cmolog /FL-qb:AB020712.1 |
| 211761_s_at | gb:BC005975.1 /DEF-Homo sapiens, calcyclin binding protein, clone MGC:14660, mRNA, complete cds. /FEA-mRNA /PROD-calcyclin binding protein /DB XREF-gi:13543650 /FL-gb:BC005975.1 |
| 213164_at | Consensus includes gb:AI867198 /FEA=EST /DB XREF=91:5540214 /DB XREF=est:wa01c11.x1 /CLONE=IAAGE:2296820 /UG=Hs.324787 solute carrier fam- 11y 5 (inositol transporters), member 3 /FL=qb:NM 006933.1 |
| 219940_s_at | gb:NM 018386.1 /DEF=Homo saplens hypothetical protein FLJ11305 (FLJ11305), mRNA. /FEA=mRNA /GEN=FLJ11305 /PROD=hypothetical protein FLJ11305 /FLJ11305 /DB XREF=gi:8922986 /UG=Hs.7049 hypothetical protein FLJ11305 /FL=ab:NM 018386.1 |
| 203800_s_at | 3 /FEA=EST /DB_XREF=g1:12764469 /DB_XREF=est:6023686 |
| 213026_at | Consensus includes gb:BE965998 /FEA-EST /DB_XREF-gi:11770950 /DB_XREF-est:601659892R1 /CLONE-IMAGE:3905710 /UG-Hs.264482 Apg12 (autophagy 12, S. cerevisiae)-like |
| 215165_x_at | Consensus includes gb:AL080099.1 / DEF=Homo sapiens mRNA; DNA DKFZp564G1272 (from clone DKF2p564G1272); partial cds. /FEA=mRNA /GEN=DKF2p564G1272 /PROD=hypothetical protein /DB_XREF=gi:5262522 /UG=Hs.2057 uridine monophosphate synthetase (orotate phosphoribosyl transferase and orotidine-5-decarboxylase) |
| 202864_s_at | gb:NM 003113.1 /DEF-Homo sapiens nuclear antigen Sp100 (SP100), mRNA. /FER-mRNA /GEN-SP100 /PROD-nuclear antigen Sp100 /DB_XREF-g1:4507164 /UG-Hs.77617 nuclear antigen Sp100 /EL-dp:M60618.1 dp:NM 003113.1 |
| 208407_s_at | iens ca), delt |
| 202419_at | gb:NM_002035.1 /DEF-Homo sapiens follicular lymphoma variant translocation 1 (FV71), mRNA. /FEA-mRNA /GEN-FV71 /FROD-follicular lymphoma variant translocation 1 /DE XREF-gi:4503016 /UG-Hs.74050 follicular lymphoma variant translocation 1 /FI-cb:NM 007035.1 |
| 201833_at. | ns histone deacetylase 2 (HDAC2), mRNA. /FEA=mRNA /GEN=HDAC2 /PROD=histc e 2 /FL=gb:U31814.1 qb:NN 001527.1 |
| 217879_at | Consensus includes gb:AL566824 /FEA-EST /DB_XREF-gi:12919571 /DB_XREF-est:AL566824 /CLONE-CSODF025YN03 (3 prime) /UG-Hs.172405 cell divi- sion cycle 27 /FL-gb:NM 001256.1 |

Table 5: Genes from Cluster Analysis 5

| Affymetrix | Description of the sequence in the GeneBank data base |
|-------------|---|
| internal | |
| designation | |
| 201466_s_at | gb:NM 002228.2 /DEE-Homo sapiens v-jun avian sarcoma virus 17 oncogene homolog (JUN), mRNA. /FEA-mRNA /GEN-JUN /PROD-v-jun avian sarcoma virus 17 oncogene homolog /DE XREF-gi:7710122 /UG-Hs.78465 v-jun avian sarcoma virus 17 oncogene homolog /FL-gb:BC002646.1 gb:NM 002228.2 |
| 202425_x_at | pb:NM 000944.1 /DEE-Homo sapiens protein phosphatase 3 (formerly 2B), catalytic subunit, alpha isoform (calcineurin A alpha) (PPP3CA), mRNA. //FEA-mRNA /GEN-PPP3CA /PROD-protein phosphatase 3 (formerly 2B), catalyticsubunit, alpha isoform (calcineurin A alpha) /DB XREF-qi:6715567 / uG-Hs:272458 protein phosphatase 3 (formerly 2B), catalytic subunit, alpha isoform (calcineurin A alpha) /FL-qb:305480.1 gb:L14778.1 hh:NN 000944.1 db:A135350.1 |
| 218566_s_at | gb:NM 012124.1 /DEF-Homo sapiens chord domain-containing protein 1 (CHPI), mRNA. /FEA-mRNA /GEN-CHPI /PROD-chord domain-containing protein 1 /PB XREF-gi:6912303 /UG-Hs.22857 cysteine and histidine-rich domain (CHORD)-containing, zinc-binding protein 1 /FL-gb:AF192466.1 dp:NH 012124.1 |
| 214349_at | Consensus includes gb:AV764378 /FEA-EST /DB XREF-g1:10922226 /DB_XREF-est:AV764378 /CLONE-MDSAOE03 /UG-Hs.163863 ESTs, Moderately similar to POLZ HUMAN RETROVIRUS-RELATED POL POLYPROTEIN H.sapiens |
| 204314_s_at | db:NM 004379.1 /DEF-Homo sapiens cAMP responsive element binding protein 1 (CREB1), mRNA. /FEA-mRNA /GEN-CREB1 /PROD-cAMP responsive elemen binding protein 1 /DB XREF-g1:4759053 /UG-Hs.79194 cAMP responsive element binding protein 1 /FL-gb:N27691.1 gb:NM 004379.1 |
| 208753_s_at | gb:BC002387.1 /DEF-Nomo sapiens, nucleosome assembly protein 1-like 1, clone MGC:8688, mRNA, complete cds. /FEA-mRNA /PROD-nucleosome assem bly protein 1-like 1 /DB XREF-gi:12803166 /UG-Hs.179662 nucleosome assembly protein 1-like 1 /FL-gb:BC002387.1 gb:AL162068.1 |
| 215452_x_at | Consensus includes gb:AL031133 /DEF-Numan DNP sequence from clone 281H8 on chromosome 6q25.1-25.3. Contains up to four novel genes, one with similarity to KIAA0323 and worm C30F12.1 and another with Ubjouitin-Like protein gene SMT3 (the latter in an intron of a novel gene). Con- |
| | tains /FEA-mRNA 2 /DB XREF-g1:3676189 /UG-Hs.113293 Human DNA sequence from clone 281H8 on chromosome 6q25.1-25.3. Contains up to four novel manes one with similarity to KTAA0323 and worm C30F12.1 and another with Ubiquitin-Like process one SMT3 (the latter in an intron of a |
| | novel gene). Contains ESTs, S |
| 204119_s_at | gb:U90339.1 /DEF=Human adenosine kinase short form mRNA, complete cds. /FEA-mRNA /PROD=adenosine kinase short form /DB_XREF=gi:1906010 / UG-Hs.94382 adenosine kinase /FL-gb:U50196.1 gb:BC003568.1 gb:U90339.1 gb:NM 001123.1 |
| 201304_at | gb:NM 005000.2 /DEF=Homo sapiens NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 5 (13kD, B13) (NDUFA5), nuclear gene encoding mitochondrial protein, mRNA. /FEA-mRNA /GEN=NDUFA5 /PROD=NADH dehydrogenase (ubiquinone) 1 alphasubcomplex, 5 /DB XREF=gi:13699821 / UG-Hs.83916 NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 5 (13kD, B13) /FL-gb:BC000813.1 gb:NM 005000.2 gb:U53468.1 gb:U64028.1 |
| 218131_s_at | gb:NM 017660.1 /DEF-Homo sapiens hypothetical protein FLJ20085 (FLJ20085), mRNA. /FEA-mRNA /GEN-FLJ20085 /PROD-hypothetical protein FLJ20085 /DB XREF-g1:8923093 /UG-Hs.118964 hypothetical protein FLJ20085 /FL-gb:NM 017660.1 |
| 201788_at | gb:NN 007372.1 /DEF-Homo sapiens RNA helicase-related protein (RNAMP), mRNA. /FEA-mRNA /GEN-RNAMP /PROD-RNA helicase-related protein / DB XREF-gi:11321631 /UG-Hs.0765 RNA helicase-related protein /FL-gb:NM 007372.1 gb:AF003255.1 |
| 209780_at | gb:AL136883.1 /DEF-Homo sapiens mRNA; cDNA DKFZp434D166 (from clone DKFZp434D166); complete cds. /FEA-mRNA /GEN-DKFZp434D166 / PROD-hypothetical protein /DB XREF-gi:12053266 /UG-Hs.128653 hypothetical protein DKFZp564F013 /FL-gb:AL136883.1 |
| 219375_at | gb:NH 006090.1 /DEE-Homo sapiens cholineethanolaminephosphotransferase (CEPT1), mRNA. /FEA-mRNA /GEN-CEPT1 / PROD-cholineethanolaminephosphotransferase /DB_XREE-gi:5174414 /UG-Hs.125031 cholineethanolaminephosphotransferase / FL-gb:AF068302.1 gb:NM 006090.1 |
| 209187_at | Consensus includes gb:AW\$16932 /FEA=EST /DB XREF=gi:7154941 /DB XREF-est:xq04a05.x1 /CLONE=IMAGE:2748848 /UG=Hs.16697 down-regulator of transcription 1, TBP-binding (negative cofactor 2) /FL=gb:BC002809.1 |
| 219279_at | gb:NM 017718.1 /DEF-Homo sapiens hypothetical protein FLJ20220 (FLJ20220), mRNA. /FEA-mRNA /GEN-FLJ20220 /PROD-hypothetical protein FLJ20220 /DB XREF-gi:8923209 /UG-Hs.21126 hypothetical protein FLJ20220 /FL-gb:NM 017718.1 |
| 206976_s_at | |
| 215596_s_at | Consensus includes gb:Al163248 /DEE-Homo sapiens chromosome 21 segment HS21CO48 /FEA-mRNA_2 /DB_XREF-g1:7717304 /UG-Hs.288773 zinc finger protein 294 |
| 216511_s_at | Consensus includes gb:AJZ70770 /DEF=Homo sapiens partial TCF-4 gene for T-cell transcription factor-4, exon 1 and joined CDS features / FER=mRNA 1 /DB XREF=gi:9188625 /UG=Hs.283857 Homo sapiens partial TCF-4 gene for T-cell transcription factor-4, exon 1 and joined CDS fea- |
| | |

| ÷ | tures |
|-------------------|---|
| 201889_at | gb:NM 014888.1 /DEF=Homo sapiens predicted osteoblast protein (GS3786), mRNA. /FEA-mRNA /GEN-GS3786 /PROD-predicted osteoblast protein / DB XREF=gi:1661713 /UG-Hs.29882 predicted osteoblast protein / FI-ch:NA 014888 |
| 201435_s_at | /FEA=EST /DB XREF=gi:6655670 /DB_XREF=est:xv52 3353.1 gb:NM 001968.1 |
| 212232_at | Consensus includes gb:AB023231.1 /DEF-Homo sapiens mRNA for KIAA1014 protein, partial cds. /FEA-mRNA /GEN-KIAA1014 /PROD-KIAA1014 protein / DB XREF-g1:4589677 /UG-Hs.6834 KIAA1014 protein |
| 214329_x_at | |
| 212099 <u>a</u> t | Consensus includes gb:AI263909 /FEA=EST /DB_XREF=g1:3872112 /DB_XREF=est:q108f09.x1 /CLONE=IMAGE:1855913 /UG=Hs.204354 ras homolog gene family, member B /FL-gb:NM 004040.1 |
| 213618_at | Consensus includes gb:AB011152.1 /DEF-Homo sapiens mRNA for KIAA0580 protein, partial cds. /FEA-mRNA /GEN=KIAA0580 /PROD=KIAA0580 protein / DB XREF-gi:3043683 /UG-Hs.22572 KIAA0580 protein |
| 216899_s_at | Consensus includes gb:AC003999 /DEF=Human PAC clone RPS-1139P1 from 7p15-p21 /FEA=CDS /DB_XREF=gi:2772566 /UG=Hs.52644 SKAP55 homologue |
| 201437_s_at | is eukaryotic translation initiation factor 4E (EIE4E), mRNA. /FEA-mRNA /GEN-EIE4E |
| 212122_at | LILLACION TACCOL 4E 705 AND TOTAL TOTAL TOTAL TOTAL TERRELATION INITIATION FACTOR 4E 7EF-95:M15353.1 95:NM 001968.1 CONSENSUS INCLUSES 95:AW711590 /FEA-EST 708 XREF-91:7703647 / DB XREF-95:1:05080.X1 / CLONE-IWAGE:3032126 / UG-Hs.166982 phosphatidylinositol q1ycan, class I |
| 208666_s_at | udes gb: BE866412 |
| 218252_at | F-Homo sapiens cytoskeleton associated protein |
| | |
| 216384_x_at | Consensus includes gb: AF257099 / DEF=Homo sapiens prothymosin alpha (PTMA) gene, complete cds /FEA=CDS / DB_XREF-gi: 8037944 / UG=Hs. 283947 Homo sapiens prothymosin alpha (PTMA) gene, complete cds |
| 217100_s_at | 1 /DEF= |
| 204313_s_at | |
| 214257_s_at | Consensus includes gb:AA890010 /FEA-EST /DB_XREF-gi:3016889 /DB_XREF-est:aj89h08.sl /CLONE-IMAGE:1403679 /UG-Hs.50785 SEC22, vesicle traf- ficking |
| | |
| 218595_s_at | |
| 213128_s_at | /FEA=EST /DB_XREF=gi:2269568 /DB_661043) |
| 207719_x_at | gb:NM 014812.1 /DEF-Homo sapiens KIAA0470 gene product (KIAA0470), mRNA. /FEA-mRNA /GEN-KIAA0470 /PROD-KARP-1-binding protein / DB XREF-gi:1662141 /UG-Hs.25132 KIAA0470 gene product /FL-qb:AB022657.1 ab:NM 014812.1 |
| 200060_s_at | IZ W |
| 200634_at | gb:N3 005022.1 /DEF-Homo sapiens profilin 1 (PFN1), mRNA. /FEA-mRNA /GEN-PFN1 /PROD-profilin 1 /DB_XREF-g1:4826897 /UG-Hs.75721 profilin 1 / FL-qb:BC002475.1 qb:303191.1 qb:NN 005022.1 |
| 202770_s_at | gb:NN 004354.1 /DEF-Homo sapiens cyclin G2 (CCNG2), mRNA. /FEA-mRNA /GEN-CCNG2 /PROD-cyclin G2 /DB_XREF-gi:4757935 /UG-Hs. 79069 cyclin G2 / FL-gb:U47414.1 gb:NN 004354.1 |
| 201008_s_at | Consensus includes gb:AA812232 /FEA-EST /DB XREF-gi:2881843 /DB XREF-est:ob84h09.s1 /CLONE-IMAGE:1338113 /UG-Hs.179526 upregulated by 1,25-dihydroxyvitamin D-3 /FL-gb:NM 006472.1 gb:S73591.1 |
| 218534_s_at | gb:NM 018046.1 /DEF-Homo sapiens hypothetical protein FLJ10283 (FLJ10283), mRNN. /FEA-mRNA /GEN-FLJ10283 /FROD-hypothetical protein FLJ10283 /FLJ (DB XREF-gi:8922325 /UG-HS.284216 hypothetical protein FLJ10283 /FL-gb:NN 018046.1 |
| | |

| 200993_at | Consensus includes gb:AA939270 /FEA=EST /DB_XREF=g1:3099183 /DB_XREF=est:oq31b02.s1 /CLONE=IMAGE:1587915 /UG=Ks.5151 RAN binding protein 7 / |
|-------------|--|
| 214102_at | ELYGOTATION STATES OF THE PROPERTY OF THE PROP |
| 213704_at | /FEA-EST /DB_XREF-g1:1690163 /DB_XREF-est:z116a12.s1 |
| 212367_at | 4GE: 2349 |
| 201873_s_at | gb:NM 002940.1 /DEF-Homo sapiens ATP-binding cassette, sub-family E (OABP), member 1 (ABCE), mkNA. / VEA-MKNA / VEN-ADEL / FNOD-AIR-DINGLING Cassette, sub-family E, member 1 /DB XREF-gi:4506558 /UG-Hs.12013.ATP-binding cassette, sub-family E, member 1 /FN-gi:NM 002940.1 |
| 218618_s_at | = = : |
| 203689 s at | 7 /FER=EST /DB_XREF=g1:5111325 /DB_XREF=e85C:WG83GU5.X1 /LLUNE=IMAGE:23.1001 /UG-m3:U9701 1.1 |
| 201074_at | 3 /FEA=EST /DB XREF=91:409333 /DB XREF=65C:NN16001.51 /LLONE=FRAGE:1004033 /OS-ns.17220 SHESH. gulator of chromatin, Subfamily c, member 1 / FL=9b:U66615.1 gb/NN 003074.1 |
| 200778_s_at | (REF=q1:3/4/268 (/DB ARRE=e81:9948903.X1 /CLOME=IRAGE:1/4/260 /OS=13:10303) |
| 202307_s_at | gb:NM 000593.2 /DEF-Homo sapiens ATP-binding cassette, sub-family B:(MDRIAP), member 2 (ABCB2), mkNA. /teA-mkNA /GEN-ABCB2 /FNOD-AIF-DINGLINGLINGLINGLINGLINGLINGLINGLINGLINGL |
| 213145_at | 6 /FEA-EST / DB_XREF=gi:10701941./DB_XREF=est:7g91d12.x1./CLONE=IMAGE:3313847 /UG-Hs.12460 Homo |
| 218588_s_at | en reading en reading |
| 217834_s_at | gb:NN 006372.1 /DEF=Homo. sapiens. NS1-associated protein I (NSALI), mkNA. / Fraemina. / Groundstat / Froundstat described protein I / Fraemina. / Groundstat described by NS1-associated protein I / Fraemina / Groundstat described from the finest of the finest protein in the finest of the finest protein in the finest protein in the finest of the finest protein in the finest of the finest protein in the finest protei |
| 205038_at | =g1:1353Z/34.7DE_XKER=eSE:bUZ389Z30EL./LLUNE=IRAME:4093/83 /00-ns.3949Z ZINC LINGE 006060.1 |
| 203132_at | cluding osteosarcoma) olastoma.l (including |
| 211297_s_at | EF-Human protei 242 /UG-Hs.1842 |
| 212375_at | (03 /DB_XREE=est:Al563/2/ /CLONE=CSUBBOU/ILFO (3 prime) /uc=ns.sucosa |
| 208986_at | Consensus includes gb:AL559478 /FERA-EST /DB XREF-gi:12905019 /DB XREF-est:AL559478 /CLONE-CSUDJUJIM1 (3 pilme) /UG-ms.4104 Clauseription factors 4) /FI-gi:M8032333.1 gb:NM 0032020-pelix-loop-helix transcription factors 4) /FI-gi:M8032333.1 gb:NM 0032020-pelix-loop-helix FRA42 F-1 enxyme |
| 217956_s_at | gb:NN 021204.1 /DEF=Homo sapiens E-1 enzyme (MASA), mkNA. /EkA=mkNA /EkN=hASA /EKOL=E-1 enzyme /DE_ANLL 91:1200020 /OC MOTEST / COMPANY |
| 214870 x at | DEF-Human Chromosome to EAC Clone C1190/SA-A-30971 /FEA-WARA / 100 ARCH 94-E-2013/A |
| 209852_x_at | <pre>gb:BC001423.1 /DEF=Homo sapiens, Similar to proteaseome (prosome, macropain) 28 subunit, 3, clone roc.1334, mark, comprese cus. 6 EEA=mRNA /PROD=Similar to proteaseome (prosome, macropain) 28subunit, 3 /DB_XREF=qi:12655138 /UG=Hs.152978 proteasome (prosome, macropain) activator subunit 3 (PA28 gamma; Ki) /EL=gb:BC001423.1.gb:BC002684.1</pre> |
| 209063_x_at | 5 /FEA=EST /DB XREF=91:11164389 /DB XREF=est:0 rotein 1 /FL=9D:AL136920.1 |
| 201177_s_at | gb:NN 005499.1 /DEF=Homo sapiens SUNO-1 activating enzyme Subunit 2 /FI=gb:BC003153.1 gb:NF090364.1 gb:AF079566.1 gb:AF079566.1 gb:AF079566.1 gb:AF079566.1 gb:AF079566.1 gb:AF079566.1 gb:AF079566.1 gb:AF10957.1 gb:AF07965.1 gb:AF10957.1 gb:AF10957.1 gb:AF078 KTABA056 Tree and AF10957.1 gb:AF10957.1 gb:AF10957.1 gb:AF10957.1 gb:AF10957.1 gb:AF10957.1 gb:AF10957.1 gb:AF10957.1 gb:AF1078 KTABA056 AF1078 AF10957.1 gb:AF10957.1 gb:AF10957.1 gb:AF10957.1 gb:AF1078 KTABA056 AF1078 AF10957.1 gb:AF10957.1 gb:AF10957.1 gb:AF1078 KTABA056 AF1078 AF10957.1 gb:AF10957.1 |
| 212264_s_at | Consensus includes gb:BE645850 /FEA=EST /DB_XREF=q1:99/Ulb1 /DB_XREF=est:/e//CU3.xi /CLONE=INAGE:5288484 /OG-U3:12:20 NICOLA PLOCETI |

| 218878_s_at | s sirtuin (silent mating type information regulation 2, S. cerevisiae, in 1 /DB XREF-gi:13775598 /UG-Hs.31176 sirtuin (silent mating type inf b:AF083106.2 |
|-------------|---|
| 202163_s_at | s CCR4-NOT transcription complex, subunit 8 (CNOT8), 4758945 /UG-Hs.26703 CCR4-NOT transcription complex, |
| 203584_at | s KIAA0103 gene product 7 KIAA0103 gene product |
| 201901_s_at | _ |
| 213743_at | Consensus includes gb:BE674119 /FEA=EST /DB_XREF=gi:10034660 /DB_XREF=est:7d75b03.xl /CLONE=INAGE:3278765 /UG=Hs.155478 cyclin T2 |
| 202883_s_at | FEA=EST /DB XREF=qi:698093 /DB XREF=est:yd71a11.s1 /CL nit A (PR 65), beta isoform /F $\overline{ m L}$ =gb:NM 002716.1 gb:AF16 |
| 202069_s_at | டிய |
| 222303_at | Consensus includes gb:AV700891 /FEA=EST /DB_XREF=g1:10302862 /DB_XREF=est:AV700891 /CLONE=CKCEQD03 /UG-Hs.292477 ESTs |
| 203102_s_at | 1 C O 1 |
| 212982_at | Consensus includes gb.A1621223 /FEA-EST /DB_XREF=g1:4630349 /DB_XREF=est:ts17a09.x1 /CLONE=IMAGE:2237272 /UG-Hs.4014 KIAA0946 protein; Huntingtin interacting protein H |
| 200050_at | s zinc finger p 9 zinc finger p |
| 202430_s_at | gp:NM 021105.1 /DEE=Homo sapiens phospholipid scramblase 1 (PLSCR1), mRNA. /FEA=mRNA /GEN=PLSCR1 /PROD=phospholipid scramblase 1 / DB XREF=gi:10863876 /UG=Hs.198282 phospholipid scramblase 1 /FL=gb:NM 021105.1 gb:AB006746.1 gb:AF098642.1 |
| 218757_s_at | ns similar /UG=Hs.10 |
| 214030_at | /FEA-EST /DB_XREF-gi:9703760 /DB_XREF-est:7a41e05.xl /CLONE-IMAGE:3221312 /UG-Hs.23294 ESTs, T28F2.4 - Caenorhabditis elegans C.elegans |
| 218093_s_at | is hypothe 372 hypoth |
| 203486_s_at | /FEA=EST /DB_XREF=gi:11083411 /DB_XREF=est:7088c12.x1 /CLONE=INAGE;3643391 /UG=Hs.102708 DKF? |
| 219303_at | .s hypothetical protein FLJ13449 (FLJ13449), mRNA. /FEA-mRNA /GEN-FLJ13449 / UG-Hs.10711 hypothetical protein FLJ13449 /FL-gb:AL136651.1 gb:NM 024546.1 |
| 202173_s_at | ns zinc finger protein 161 (ZNF161), mRNA. /FEA-mRNA /GEN-ZNF161 /PROD-zinc finger zinc finger protein 161 /FL-gb:D28118.1 gb:NM 007146.1 |
| 216903_s_at | 1 /DEF=Homo sapiens cDNA FLJ12635 fis, clone NT2RM -mRNA /DB XREF=gi:10434244 /UG=Hs.61628 calcium bi |
| 205842_s_at | s Jak2 kinase (JAK2) mRNA, c protein tyrosine kinase) /E |
| 212702_s_at | EA-ESI /DB XREF-gi:1186277 /DB XREF-est:yz12f12.s1 /CLONE-INAGE:282 11, clone IMAGE:3622452, mRNA, partial cds |
| 201664_at | DEF»Homo sapiens mRNA; cDNA DKF2p434F205 (from clone DKF2p434F205); complete cds. /FEA»mRNA /GEN»DKF2 cal protein /DB_XREF»gi:6807670 /UG»Hs.50758 SMC4 (structural maintenance of chromosomes 4, yeast)-lil gb:AL136877.1 |
| . 202060_at | s KIAA0155 gene product (KIAA0 8 KIAA0155 gene product /FL-gb |
| 203177_x_at | gb:NM 003201.1 /DEF-Homo sapiens transcription factor 6-like 1 (mitochondrial transcription factor 1-like) (TCF6L1), mRNA. / EFE-mRNA /EEN-TCF6L1 /PROD-transcription factor 6-like 1 (mitochondrialtranscription factor 1-like) /DB XREF-giis601400 / Inclusion of Factor 6-like 1 (mitochondrial transcription factor 1-like) /Filarh.MC510 1 /DE XREF-giis601400 / |

| 209748_at | is mRNA for KIAA1083 protein, complete cds. /FEA=mRNA is 4 (autosomal dominant; spastin) /FL=qb:AB029006.1 |
|--------------|--|
| 200660_at | /Ude#13.20334 Spacert parprogres 100 calcium-binding protein All (calgizzarin) (SIOOALI), mRNA. /EEA-mRNA /GEN=SIOOALI /PROD=SIOO gb:NN 005620.1 /DEF=Homo sapiens SIOO calcium-binding protein All (calgizzarin) /FL=gb:D49355.1 calcium-binding protein All (calgizzarin) /FL=gb:D49355.1 |
| 222119_s_at | 0.1 DEF-Homo sapiens mRNA; cDNA DKF25564K2364 (From clone DKF25564K2364); partial c hetical protein /DB XREF-g1:5912200 /UG-Hs 284289 vitiligo-associated protein VIT-1 |
| 208934_s_at | ns colorectal carcinoma-derived galectin-8 variant I mRNA, complete cds. /FEA variantI /DB XREF-gi:13249298 /UG-Hs.4082 lectin, galactoside-binding, soluble qb:AF074000.I gb:NM 006499.1 |
| 211698_at | gb:AF34944.1 /DEE=Homo sapiens Rb- and p300-binding protein EID-1 mRNA, complete cds. /FEA=mRNA /PROD=Rb- and p300-binding protein EID-1 / DB XREF=qi:13549113 /FI=gb:AF9444.1 |
| 212053_at | Consensus includes gb:AKO25504.1 /DEF-Homo sapiens CDNA: FLJ21851 fis, clone HEFUL962. /FLA-mKNA /UB AREE gb:IU43041 /UG-n3.170210 ALANGEST DEOCETH |
| 209174_s_at | hypothetical protein FLJ20259, clone MC:3396, mkNA, complete cas. |
| 203476_at | gp:NN 006670.1 /DEE=Homo sapiens 5T4 oncofetal trophoblast glycoprotein (FLA=mink, /GEA=1, /FNU;-314 UNCLECEL LICPHOBLAST GLY) (MG-HS. 82128 5T4 Oncofetal trophoblast glycoprotein /FDEPSH MARKEF=41:5729717 /UG-HS. 82128 5T4 Oncofetal trophoblast glycoprotein /DB XREF=q1:5729717 /UG-HS. 82128 5T4 Oncofetal trophoblast glycoprotein /DB XREF=q1:5729717 /UG-HS. 82128 5T4 Oncofetal trophoblast glycoprotein /DB XREF=q1:4674009 /UG-HS. 12540 |
| 203007_x_at | JD:AF077198.1 /DEF=Homo sapiens lysophospholipase mkNA, Complete cds. /FrA-minna / FROU-1/sopholipase I /FL-gp:AF081281.1 gb:AF077198.1 gb:AF07128.1 gb:AF077198.1 gb:AF07 |
| 212721_at | 0 /FEA-EST /DB XKEF-91:0390940 /DB AKEF-95-1:WOO /ILC.AI / CEONELINGS.ESTE / CO. |
| 218461_at | ens protein x 0004 (LOC31184), mkna. / FEATHRING (GENT-LOCA1134 / LINE FACCOLI CO. / J. T. J. OB: NM 016301.1 |
| 214218_s_at | 7 /FEA=EST /DB_XREF=91:10301318 /DB_2 |
| 218178_s_at | db:NM 020412.1 DEF=Homo sapiens CHMP1.5 protein (CHMP1.5), mkNA. / FEA=mkNA / GEN=CHMP1.5 / FROD-CHMP1.5 protein / DEF=Homo sapiens CHMP1.5 protein / CHMP1.5 protein / FI=nh:AF281064 1 db:NM 020412.1 |
| 217851_s_at | -miring 1. PROD-CGI-107 protein (LOC51012), mRNA. /FEA-mRNA /GEN-LOC51012 /PROD-CGI-107 protein /DB_XREF-gi:7705609 / GD-104 016045.1 /DEF-Homo sapiens CGI-107 protein /FL-qb:AF151865.1 gb:NM 016045.1 |
| 202329_at | ens c-src tyrosine kinase (CSK) nase /FL=qb:NM 004383.1 |
| 203302_at | db:NN 000788.1 /DEF-Homo sapiens deoxycytidine kinase (DCK), mRNA. /FEA-mRNA./GEN=UCK /FKUD=deoxycytidine kinase /UD-ANEI-91.300250 / UG-HS-709 deoxycytidine kinase /FL-gb:N60527.1 gb:NN 000788.1. |
| .201949_x_at | 11 /EEA-EST /DB XREF=01:12530514 /DB XREF=68t:AD3/6341 /CLONE-COULDO/1253 /5 FLAME, /CO |
| 218096_at | ens hypothetical protein LO11210 (LLD1220), MRNN. (FEA-MANN. GENELLOILLY (FEA-MANN.) GENELLOILLY (FEA-MANN.) (GENELLOILLY |
| 203255_at | gb:NM 018693.1./DEF=Homo sapiens Vili1go-associated protein VII-1/FL-gb:NM 018693.1.gb:REF=gi:10048403.7UG-Hs.284289 vitiligo-associated protein VII-1/FL-gb:NM 018693.1.gb:REF=gi:10048403.7UG-Hs.284289 vitiligo-associated protein VII-1/FL-gb:NM /CENL-THARGA DEAR-KIAAARA AFREF=gi:7662363/ |
| 203855_at | gb:NH 014969.1 /DEF-Homo sapiens KIAA0893 protein (KLAA0893), mkwa. /Era-mwa./GEN-Ara-obs.// /IND-ARA0893 protein /FL-gb:AB020700.1 gb:NM 014969.1 UG-HS-3830 KIAA0893 protein /FL-gb:AB020700.1 gb:NM 014969.1 |
| 218846_at | tens cofactor required for Spl transcriptional activation, Subdult 3 (190kb) (GASTS), MANY: (TEXTMEN) transcriptionalactivation, Subunit 3 (130kb) /DB XREF-gi:1019352 /UG-Hs.29679 cofactor required for transcriptionalactivation, Subunit 3 (130kb) /FL-gb:AF135022.1 gb:AF105332.1 gb:NM 004830.1 |
| 208988_at | 1843 /FEA=EST /D -9b:AF179221.1 |
| 216202_s_at | 1. / Ubrahuman Seline paimitoyitianistesso |
| 216954_x_at | Consensus includes gb:S//336.1 / UEr=nomo Saprens Originaria Sanstaria Consensus includes gb:S//336.1 / UEr=nomo Saprens Originaria Sanstaria Sans |
| | |

| · | nferral proteinoscp-like protein /DB_XREF=gi:944964 /U cin sensitivity conferring protein). |
|-------------|---|
| 212753_at | Consensus includes gb:AI692203 /FEA=EST /DB_XREF=g1:4969543 /DB_XREF=est:wd37h12.x1 /CLONE=IMAGE:2330375 /UG=Hs.8834 ring finger protein 3 |
| 207691_x_at | gb:NM 001776.1 /DEF-Homo sapiens ectonucleoside triphosphate diphosphohydrolase 1 (ENTPD1), mRNA. /FEA-mRNA /GEN-ENTPD1 /PROD-ectonucleosid triphosphate diphosphohydrolase 1 /FL-gb:NM 001776.1 |
| 200967_at | C CO |
| 201988_s_at | Consensus includes gb:BF438056 /FEA=EST /DB_XREF=gi:11450573 /DB_XREF=est:7q66e01.x1 /CLONE=IMAGE:3703369 /UG=Hs.13313 CAMP responsive element binding protein-like 2 /FI=gb:AF039081.1 gb:NM 001310.1 |
| 202142_at | gb:BC003090.1 /DEF-Homo sapiens, COP9 homolog, clone MGC:1297, mRNA, complete cds. /FEA=mRNA /PROD-COP9 homolog /DB_XREF=g1:13111846 / UG-Hs.75193 COP9 homolog /FL-gb:BC003090.1 gb:U51205.1 gb:NM 006710.1 |
| 218238_at. | gb:NM 012341.1 /DEF=Homo sapiens GTP-binding protein (NGB), mRNA. /FEA-mRNA /GEN=NGB /PROD=GTP-binding protein /DB_XREF=gi:6912531 / UG=Hs.215766 GTP-binding protein /FL=gb:AF325353.1 gb:AF120334.1 gb:NM 012341.1 |
| 205552_s_at | 3b:1M 002534.1 /DEF=Homo sapiens 2,5-oligoadenylate synthetase 1 (40-46 kD) (OAS1), transcript variant E16, mRNA. /FEA-mRNA /GEN-OAS1 / DRON-2 5-olicoadenylate synthetase 1 isoform E16 /DB XREF=01:8051622 /UG-Hs 82396 2 5-olicoadenylate synthetase 1 (40-46 kD) |
| - | |
| 202541_at | Consensus includes gb:EF589679 /FEA-EST /DB XREF-gi:11682003 /DB XREF-est:naa08b05.x1 /CLONE-INAGE:3253977 /UG-Hs.146401 small inducible cytokine subfamily E, member 1 (endothelial monocyte-activating) //FL-gb:NM 004757.1 gb:U10117.1 |
| 221699_s_at | gb:AF334103.1 /DEF-Homo sapiens nucleolar protein GU2 mRNA, complete cds. /EEA-mRNA /PROD-nucleolar protein GU2 /DB XREF-g1:13540303 / FL-gb:AF334103.1 |
| 202090_s_at | gb:NM 006830.1 /DEF=Homo sapiens ubiquinol-cytochrome c reductase (6.4kD). subunit (UQCR), mRNR. /FEA=mRNR /GEN=UQCR / PROD=ubiquinol-cytochrome c reductase (6.4kD)subunit /DB_XREF=gi:5803216 /UG=Hs.8372 ubiquinol-cytochrome c reductase (6.4kD) subunit / FL=db:D55636.1 db:NM:006830.1 |
| 202382_s_at | db:NM 005471.1 /DEF-Homo sapiens glucosamine-6-phosphate isomerase (GNPI), mRNA. /FEA-mRNA /GEN-GNPI /PROD-glucosamine-6-phosphate isomerase /FL-gb:NN 005471.1 gb:AF029914.1 gb:AF048826.1 gb:D31766.1 |
| 220408 x at | gb:NN 017569.1 /DEF-Homo sapiens transcription factor (p38 interacting protein) (F36IF), mRNA. /FEA-mRNA /GEN-F38IF /PROD-transcription factor (p38 interacting protein) /DB_XREF-gi:8923734 /UG-Hs.171185 transcription factor (p38 interacting protein) /FL-gb:AFC93250.1 |
| 208264_s_at | gp. na. 01705:: gb:NN 003758.1 / DEF-Homo sapiens eukaryotic translation initiation factor 3, subunit 1 (alpha, 35kD) (EIF3S1), mRNA. / FEA-mRNA / GEN-EIF3S1 PROPEUKARYOTIS I Initiation factor 3, subunit 1 (alpha, 35kD) / DB_XREF-gi:4503510 / UG-Hs.173987 eukaryotic translation initiation factor 25kD) - Learn Angelon 1 (alpha, 35kD) / DB_XREF-gi:4503510 / UG-Hs.173987 eukaryotic translation initiation factor 3 (alpha, 35kD) - Learn Angelon 1 (alpha, 35kD) / DB_XREF-gi:4503510 / UG-Hs.173987 eukaryotic translation initiation |
| 202872_at | ensus includes gb:AW024925 /FE |
| | p) 42kD /FL=gb:NM 001695.1 |
| 214835_s_at | Consensus includes gb:AF131748.1 /DEF=Homo sapiens clone 25191 GTP-specific succinyl-CoA synthetase beta subunit (SCS) mRNA sequence, partial cds. /FEA=mRNA /PROD=GTP-specific succinyl-CoA synthetase betasubunit /DB_XREF=gi:4406563 /UG=Hs.247309 succinate-CoA ligase, GDP-forming, beta subunit |
| 205263_at . | gb:AF082283.1 /DEF=Homo sapiens CARD-containing apoptotic signaling protein (BCL10) mRNA, complete cds. /FEA=mRNA /GEN-BCL10 / PROD=CARD-containing apoptotic signaling protein /DB XREF=gi:4092066 /UG=Hs.193516 B-cell CLLlymphoma 10 /FL=gb:AF082283.1 ph:AF12737 |
| 218512_at | hypotheti Hs.73291 |
| 210283_x_ac | s, Similar to polyadenylate binding protelr adenylate bindingprotein-interacting protei /FL=gb:BC005295.1 |
| 202724_s_at | ens 133 |
| 212058_at | /FEA=EST /DB_XREE=gi:3735200 /DB_XREF=est:qd60b05.xl / |

| | | 1 |
|-----------|-------------|--|
| 212006_at | | Consensus includes gb:AU149908 /FEA-EST /DB_XREF=gi:11011429 /DB_XREF=est:AU149908 /CLONE=NT2RP2000896 /UG=Hs.//495 UBX |
| 222040_at | | s gb:AI144007 |
| 20237 | 102373_s.at | gb.AF755648.1 /DEE-Homo sapiens rGAP-iso mRNA, complete cds. /FEA-mRNA /PROD-rGAP-iso /DB XREF-gi:12005820 /UG-Hs.197289 rab3 GrPase-activating protein, non-catalytic subunit (150kD) /FL-gb:AF255648.1 gb:AF004828.1 gb:NN 012414.1 |
| 209301_at | 1_at | 0 21 |
| 213264_at | 14_at | œ'. L |
| 202863_at | 3_at | gb:NM 003113.1 /DEF=Homo sapiens nuclear antigen Sp100 (SP10U), mkNA: /FEA=mkNA /GEN=SP10U /FNOD=nuclear antigen Sp100 /DE_AREF=41:430:100 / UG-HS:77617 nuclear antigen Sp100 /FI=qb:M60618.1 gb:NN 003113.1 |
| 21767 | 217679_x_at | _ E |
| 21280 | 12802_s_at | Consensus includes gb:AK023841.1 / DEF-Homo sapiens cDNN FLJ3779 fis, clone PLACE40U0445, nighry similar to homo sapiens mana; conva DKF2p434C212 (from clone DKF2p434C212): /FEA-mRNA /DB XREF-gi:10435900 /UG-Hs.172069 DKF2P434C212 protein |
| 21034 | 210346_s_at | ns CLK4 mRNA, complete cds. /FEA. |
| 20884 | 108848_at | |
| 221978_at | 18_at | |
| 21752 | 217526_at | jb.aI478300 /FEA-EST /DB XREF-gi:43115.6 /DB XREF-est:tm39e01.XI /CLONE=IMAGE:ZIG0304 /UG=N3:192703 E3:2, Wear JBFAMILY SP SEQUENCE CONTAMINATION WARNING ENTRY H.sapiens |
| 2183 | 218322_s_at | gb:NM 016234.2 /DEF=Homo sapiens long-chain fatty acid coenzyme A 11gase 5 (FACL5), mANA. /FEA=mRNA /GEN=FACL5./FROU=LUNG-Chain latry acid coenzyme A 11gase 5 /DB XREP=gi:12669912 /UG=Hs:11638 long-chain fatty acid coenzyme A 11gase 5 /FLAGD-LUNG-GEN-FACL5. GEN-FACL5 (MC-Hs BAA01 hybrosphilis) |
| 2126 | 12613_at | Consensus includes gb:AI991252 /FEA-EST /DB_XKEF=91:383813/ /DB_AKEF=55:WutteU9:X1 /CLUNE-InfoGE:222202 /UG-13:0:4:4/ Date Consensus includes 3, member A2 /UG-13:0:4/ /CE11-DBM10 / |
| 2045 | 204566_at | gb:NM 003620.1 /DEF=Homo sapiens protein phosphatase 1D magnesium-dependent, detta isotorm (Proid), minkr. /refrance / Defense in magnesium-dependent, delta isoform /DB_XREF=q1:4505996 /UG=Hs.100980 protein phosphatase 1D magnesium-dependent, delta isoform /EL=qp:U78305.1 gb:NM 003620.1 |

Table 6: Genes from Cluster Analysis 6

| Affymetrix internal | Description of the sequence in the GeneBank data base |
|------------------------|---|
| designation | |
| 202018_s_at | gb:NM 002343.1 /DEF-Womo sapiens lactotransferrin (LTF), mRNA. /FEA-mRNA /GEN-LTF /FROD-lactotransferrin /DB_XREF-gi:4505042 /VG-Ws.105938 lactotransferrin /FL-gb:AF332168.1 gb:M83202.1 gb:NM 002343.1 |
| 216379_x_at | Consensus includes gb:AK000168.1 /DEF=Homo sapiens cDNA FLJ20161 fis, clone COL09252, highly similar to L33930 Homo sapiens CD24 signal transducer mRNA. /FEA=mRNA /DB XREF=gi:7020079 /UG-Ms.332045 Homo sapiens cDNA FLJ20161 fis, clone COL09252, highly similar to L33930 Homo sapiens CD24 signal transducer mRNA |
| 209771_x_at | Consensus includes gb.AA761181 /FEA=EST /DB XREF=gi:2810111 /DB XREF=est:nz09q03.s1 /CLONE=IWAGE:1287316 /UG-Hs.286124 CD24 antigen (small cell lung carcinoma cluster 4 antigen) /FL=gb:X69397.1 |
| 206676_at | gb:M33326.1 /DEF=Numan nonspecific cross-reacting antigen (NCA) mRNA, complete cds. /FEA=mRNA /GEN-MCA /PROD-non-specific cross reacting antigen / B XREF=g1:189101 /UG-Hs.41 carcinoembryonic antigen-related cell adhesion molecule 8 /FL-gb:M33326.1 gb:NM 001816.1 |
| 210254_at | gb:L35848.1 /DEF-Homo sapiens IgE receptor beta chain (HTm4) mRNA, complete cds. /FEA-mRNA /GEN-HTm4 /PROD-IgE receptor beta subunit /DB XREF-gi:561638 /UG-Hs.99960 membrane-spanning 4-domains, subfamily A, member 3 (hematopoletic cell-specific) / FL-gb:NM 006138.1 gb:L35848.1 |
| 203887_s_at | gb:NM 000361.1 /DEF=Homo sapiens thrombomodulin (THBD), mRNA. /FEA-mRNA /GEN-THBD /PROD-thrombomodulin /DB_XREF=g1:4507482 / UG-Hs.2030 thrombomodulin /FL-gb:M16552.1 gb:NM 000361.1 |
| 212768_s_at | Consensus includes gb:AL390736 /DEE-Human DNA sequence from clone RP11-209319 on chromosome 13 Contains ESTs, STSs and GSSs. Contains the gene for the GW112 protein with two isoforms (GW112 and KIAA4294) /FEA-mRNA /DB_XREF-gi:11102238 /UG-Hs.273321 differentially expressed in hematopoietic lineages |
| 202887_s_at | gb:NM 019058.1 /DEE=Homo saplens hypothetical protein (FLJ20500), mRNA. /FEA-mRNA /GEN=FLJ20500 /PROD=hypothetical protein / DB XREF-gi:9506686 /UG-Hs.111244 hypothetical protein /FI-gb:AL136668.1 gb:NM 019058.1 |
| 200916_at | gb:NM 003564.1 /DEF=Homo sapiens transgelin 2 (TAGLN2), mRNA. /FEA=mRNA /GEN-TAGLN2 /PROD-transgelin 2 /DB_XREF=gi:4507356 / UG-Hs-75725 transgelin 2 /FL-gb:D21261.1 gb:NM 003564.1 |
| 206157_at | ကြောင်း |
| 209369_at | gb:M63310.1 /DEF-Human 1,2-cyclic-inositol-phosphate phosphodiesterase (ANX3) mRNA, complete cds. /FEA-mRNA /GEN-ANX3 / PROD-1,2-cyclic-inositol-phosphate phosphodiesterase /DB_XREF-gi:178696 /UG-Hs.1378 annexin A3 /FL-gb:BC000871.1 gb:M63310.1 gb:M20560.1 gb:MM 005139.1 |
| 202497_x_at | Consensus includes gb:AI631159 /EEA-EST /DB XREF-g1:4682489 /DB XREF-est:ts93d05.x1 /CLONE-INAGE:2238825 /UG-Hs.7594 solute carrier family 2 (facilitated glucose transporter), member 3 /FL-qb:M20681.1 qb:NM 006931.1 |
| 205513_at | gb:NN 001062.1 /DEF-Homo sapiens transcobalamin I (vitamin B12 binding protein, R binder family) (TCN1), mRNA. /FEA-mRNA / GEN-TCN1 /PROD-transcobalamin I (vitamin B12 binding protein, Rbinder family) /DB XREF-gi:4507406 /UG-Hs.2012 transcobalamin I (vitamin B12 binding protein, R binder family) /FL-gb:J05068.1 gb:NM 001062.1 |
| 206697_s_at | gb:NM 005143.1 /DEF-Homo sapiens haptoglobin (HP), mRNNA. /FEA-mRNA /GEN-HP /PROD-haptoglobin /DB_XREF-gI:4826761 /UG-Hs.75990 haptoglobin /FL-gb:K00422.1 gb:L29394.1 gb:NM 005143.1 |
| 202146_at | Consensus includes gb:AA74726 /FEA-EST /DB XREF-gi:2787384 /DB XREF-est:nx88e08.s1 /CLONE-IMAGE:1269350 /UG-Hs.7879 interferon- related developmental regulator 1 /FL-gb:BC001272.1 gb:NM 001550.1 |
| 266_s_at | L33930 /FEATURE / DEFINITION-HUMCD24B Homo sapiens CD24 signal transducer mRNA, complete cds and 3 region |
| 211657_at | gb:H18728.1 /DEF=Human nonspecific crossreacting antigen mRNA, complete cds. /FEA=mRNA /GEN=NCA; NCA; NCA /PROD=non-specific cross reacting antigen /DB XREF=gi:189084 /FL=gb:M18728.1 |
| 205214_at | gb:NN 004226.1 /DEF-Homo sapiens serinethreonine kinase 17b (apoptosis-inducing) (STK17B), mRNA. /FEA-mRNA /GEN-STK17B / PROD-serinethreonine kinase 17b(apoptosis-inducing) /DB XREF-gi:4758193 /UG-Hs.120996 serinethreonine kinase 17b |
| | |

| | (apoptosis-inducing) /FL-gb:AB011421.1 gb:NM_004226.1 |
|-------------|--|
| 207574_s_at | gb:NM 015675.1 /DEF-Homo sapiens growth arrest and DNA-damage-inducible, beta (GADD45B), mRNA. /FEA-mRNA /GEN~GADD45B / prop-DKF2P566B133 protein /DB_XREF-gi:9945331 /UG=Hs.110571 growth arrest and DNA-damage-inducible, beta /FL-gb:AF090950.1 nb:NN 015675.1 |
| 206343_s_at | gb:NN 013959.1 /DEF=Homo sapiens neuregulin 1 (NRG1), transcript variant SMDF, mRNA. /FEA=mRNA /GEN=NRC1 /PROD=neuregulin 1 isoform SNDF //DB XREE=gi:7669517 /UG=Hs.172816 neuregulin 1 /FL=gp:NM 013959.1 gb:L41827.1 |
| 214146_s_at | /FEA=EST /DB_XREF=gi:836009 /DB_XRE Ic protein (includes platelet basic |
| 215716 s at | connective tissue-activating peptide III, neutrophil-activating peptide-1/ Consensus includes gb:L14561 /DEF-Homo sapiens plasma membrane calcium ATPase isoform 1 (ATP281) gene, alternative |
| | splice products, partial cds /FEA-mRNN 2 /DB XREF-gi:4165324 /UG-Hs.78546 ATPase, Ca++ transporting, plasma memorane 1 |
| 201179_s_at | gb:00300:1 / DEFNUMBA ALERINALYS GENERAL CONTROLLY FUNDED FOR THE PROPERTY (G Protein), compiled binding protein (G protein), compiled binding protein (G protein), alpha inhibiting Ferivity polypeptide. 3 / FEL-gb:003005:1 gb:M27543.1 gb:M27543.1 gb:NN 006496.1 |
| 201626_at | T /DB XREF-gi:13050848 /DB XREF-est:602386668F1 lin induced gene l'/FL-gb:NM 005542.1 |
| 209728_at | gb:BC005312.1 /DEF=Homo sapiens, clone MGC:12387, mRNA, complete cds. /FEA=mRNA /FROD=Unknown (protein for MGC:12387) / DB XREF=gi:13529055 /UG-Hs.318720 Homo sapiens, clone MGC:12387, mRNA, complete cds /FL=gb:BC005312.1 gb:M16942.1 |
| 212665_at | NB_XREF=g1:12899113 / DB_XREF=est:AL558438 /CLONE=CSUDKOUGILLS |
| 200665_s_at | pien otei /FL |
| 201407_s_at | .X1 /CLONE=IRAGE |
| , 203505_at | gb:AF285167.1 /DEF=Homo sapiens ATP-binding cassette transporter 1 (ABCA1) mRNA, complete cds. /FEA=mRNA /GEN=ABCA1 /PROD=ATP-binding cassette transporter 1 /DB XREF=gi:9755158 /UG=Hs.211562 ATP-binding cassette, sub-family A. (ABC1), member 1 /FL=gb:AF165281.1 gb:NM 005502.1 gb:AF285167.1 |
| 221802_s_at | Consensus includes gb:AU157109 /FEA-EST /DB_XREF-gi:11018630 /DB_XREF-est:AU157109 /CLONE-FLACE1006159 /VG-Hs.23740 KIAA1598 protein |
| 201844_s_at | <pre>2 /FEA=EST /DB XREF=g1:1395613 /DB XREF=est:zdB binding protein /FL=gb:AF179286:1.gb:AB029551.1</pre> |
| 206390_x_at | gb:NM 002619.1 /DEF=Homo sapiens platelet factor 4 (FP4); mRNA: /FEA=mRNA /GEN=F4 /FROD=platelet factor 4 /FL=gb:NJ5897.1 gb:NM 002619.1 //OB XREF=gi:4505732 //OG=H3.81564 platelet factor 4 /FL=gb:NJ5897.1 gb:NM 002619.1 |
| 212569_at | Consensus includes gb:AV699744 /FEA=EST /DB_XREF=g1:10301715 /DB_XREF=est:AV699744 /CLUNE=GRCEDEUS /UG=RS.0110 NIAAV0330 PIOCEIN |
| 211960_s_at | 416 /FEA=EST / DB XREF=g1:12 / 11.22 / DB KREF=eST:00.23/319281 .237955 hypothetical protein PRO2706 |
| 202422_s_at | gb:NM 022977.1 /DEE=Homo sapiens fatty-acid-Coenzyme A 11gase, long-chain 4 (FACL4), transcript vaidant 2, many. /FER=mRNA /GEN=FACL4 /PROD=long-chain fatty-acid-Coenzyme A 11gase 4,isoform 2 /DB XREF=gi:12669908 /UG-Hs.81452 fatty-acid-Coenzyme A 11gase, long-chain 4 /FL=gb:NM 022977.1 |
| 221958_s_at | Consensus includes gb:AA775681 /FEA=EST /DB_XREF=g1:2855015 /DB_XREF=est:z131aUz.s1 /LLURE=INAGE:3/0500 /UG=Hs.250746 hypothetical protein FLJ23091 |
| 204621_s_at | Consensus includes gb:AI935096 /FEA=EST /DB_XREF=g1:5673966 /DB_XREF=est:Wpi3el0.xl /CLONE=IM4GE:2404/40 /UG-HS.82120 nuclear receptor subfamily 4, group A, member 2 /FI—gb:NM 006186.1 |
| 201409_s_at | gb:NN 002709.1 /DEE=Homo sapiens protein phosphatase 1, catalytic subunit, Deta isoloim (FFF1cb), mann. /FEA=mRNA /GEN=PPPICB /PROD=protein phosphatase 1, catalytic subunit, betaisoform /DB XREF=gi:4506004 /VG=Hs.21537 protein phosphatase 1, catalytic subunit, beta isoform /FL=gb:NN 002709.1 gb:AF092905.1 |
| 201662_s_at | gb:DB9053.1 /DEF=Homo sapiens mRNA for Acyl-CoA synthetase 3, complete cds. /FLM=mRNA /FRUJ=Acyl-CoA synthetase 3 /DB XREF=gi:4165017 /UG=Hs.268012 fatty-acid-Coenzyme A ligase. long-chain 3 /Fl=gb:mN 004457.2 gp:DB9053.1 gb:AF116690.1 |
| 220987_s_at | gb:NM 030952.1 /DEF=Homo sapiens hypothetical protein DKFZp434J037 /DB XREF=qi:13569921 /FL-qb:NM 030952.1 |
| | |

| 219607_s_at | membrane-s ne-spannin -domains, |
|-------------------|---|
| 201565_s_at | iens inhibitor of DNA binding 2, dominant negative heli; NA binding 2, dominant negativehelix-loop-helix protein NA binding 2, dominant negative helix-loop-helix proteir |
| 213327_s_at | 101 /FEA=EST /DB XREF=gi:5439180 /DB XREF=est:wq56d12.x1 42400 ubiquitin specific protease 12 |
| 203603_s_at | gb:NM 014795.1 /DEF=Homo sapiens zinc finger homeobox 1B (ZFHX1B), mRNA. /FEA=mRNA /GEN=ZFHX1B /FROD=zinc finger homeobox 1B / FE=gb:AB011141.1 gb:NM 014795.1 |
| 217963_s_at | gb:NW 014380.1 /DEF=Homo sapiens p75NTR-associated cell death executor; ovarian granulosa cell protein (13kD) (DXS6984E), mRNA. / FEA-mRNA /GEN-DXS6984E /PROD-p75NTR-associated cell death executor; ovariangranulosa cell protein (13kD) /DB XREF=g1:1657043 / UG-Hs.1775 p75NTR-associated cell death executor; ovarian granulosa cell protein (13kD) /FI=ob:NM 014380.1 ob:AF187064 1 |
| 206710_s_at | iens differentially expressed in adenocarcinoma of the lung (KIAA0987), mRNA. (OD-differentially expressed in adenocarcinoma of the lung /DB XREF-gi:6912469 mbbane protein band 4 1~11kg 3 /FIFFment afth AD00771 ph.NM 012707 1 |
| 204342_at | |
| 204393_s_at | ppiens acid phosphatase, prostate (ACPP), mRNA. /FEA-mRNA /GEN-ACPP natase precursor /DB_XREF-gi:6382063 /UG-Hs.1852 acid phosphatase, prostate con 001099.2 |
| 200719_at | Consensus includes gb:BE964043 /FEA=EST /DB XREF=q1:11767371 /DB XREF=est:601657616R1 /FL=qb:NM 003197.2 // CLONE=IMAGE:3875955 /UG=Hs.171626 transcription elongation factor B (SIII), polypeptide 1-like /FL=qb:NM 003197.2 |
| 202731_at | iens programmed cell death 4 (PD 6251 programmed cell death 4 /FI |
| 204094_s_at | gb:NM 014779.1 /DEF-Homo sapiens KIAA0669 gene product. (KIAA0669), mRNA. /FEA-mRNA./GEN-KIAA0669 /PROD-KIAA0669 gene product / DB XREF-gi:1662235 /UG-Hs.52526 KIAA0669 gene product./FI-gb:AB014569.1 gb:NM 014779.1 |
| 208690_s_at | ens, Similar to LIM protein, clone NGC:5344, LIM protein /DB XREF=gi:12654194 /UG=Hs.755 |
| 208979_at | |
| 201772_at | piens antizyme inhibitor (LOC51582), mRNA. /FEA. 23014 antizyme inhibitor /FL=qb:D88674.1 qb:NM |
| 201345_s_at | lens ubiquitin-conjugati -ubiquitin-conjugating e 108332 ubiquitin-conjugat 3.1 |
| 221060_s_at | gb:NM 003266.1 /DEF=Homo sapiens tol1-like receptor 4 (TLR4), mRNA, /FEA=mRNA /GEN=TLR4 /PROD=tol1-like receptor 4 /DB XREF=q1:4507532 /UG=Hs.159239 tol1-like receptor 4 /FL=cb:UBBBBD 1 ob:NM 003266.1 |
| 208876_s_at | iens PAKZ mRNA, complete cds. /FEA-mRNA /PROD=PAKZ /DB_XREF=gi:5138913 PAKZ mRNA, 7complete cds /FL-qb:AF092132.1 |
| 212628_at | 065 /FEA=ES |
| 212928 <u>a</u> t | Consensus includes gb:ALO50331 /DEF=Human DNA sequence from clone 48613 on chromosome 6q22.1-22.3. Contains the part of a gene for a novel protein, the gene for KIAA0721 (NAP (Nucleosome Assembly Protein) Johnain containg protein), the TSPYL gene for TSPY-like (testis specific protein /FEA=mRNA_1 //DB XRRF-q1:5668655 (NG=Hs.28014) RTAA0721 protein / FLacb:NN 021648.1 |
| 221568_s_at | gb:AF090900.1 /DEF=Homo sapiens clone HQ0189 PR00189 mRNA, complete cds. /FEA=mRNA /PR0D=PR00189 /DB_XREF=g1:6690176 /UG=Hs.91393 Homo sapiens cDNA: FLJ21887 fis, clone HEP03135, highly similar to AF090900 Homo sapiens clone HQ0189 PR00189 mRNA /FL=gb:AF090900.1 |
| 201408 at | Consensus includes gb:W67887 /FEA-EST /DB_XREF-eg1:1376776 /DB_XREF-est:zd38c11.s1 /CLONE=IMAGE:342932 //UG-Hs.21537 protein phosphatase 1,catalytic subunit, beta isoform /FL-gb:NM 002709.1 gb:AF092905.1 |

| 202270_at | gb:NM 002053.1 /DEF-Homo sapiens guanylate binding protein 1, interferon-inducible, 67kD (GBP1), mRNA. /FER-MRNA /GEN-GBP1 /PROD-guanylate binding protein 1, interferon-inducible, 67kD /DB XREF-gi:4503938 Inches 67661 guanylate binding protein 1, interferon-inducible, 67kD /FI-gb:BC002666.1 gb:M5542.1 gb:NM 002053.1 |
|-------------|--|
| 202606_s_at | piens tousled-like kinase 1 (TLK) 18895 tousled-like kinase 1 /FL |
| 216037_x_at | |
| 221561_at | Nolesterol acyltransferase mRNA, complete cds. ltransferase /DB XREF=gl:4878021 yme A: cholesterol acyltransferase) 1 /FL=gb:NM 00310 |
| 203680 at | |
| 215111_s_at | Consensus includes gb:AKO27071.1 /DEF#Homo sapiens cDNA: FLJ2341B fis, clone HEK21445, highly similar to HSU35048 Human TSC-22 protein mRNA. /FEA=mRNA /DB XREF=gi:10440100 /UG-Hs.114360 transforming growth factor beta-stimulated protein TSC-22 |
| 202211_at | gb:BCG05122.1 /DEF=Homo sapiens, ADP-ribosylation factor GTPase activating protein 1, close MGC:10272, mRNA, complete cds. /FEA-mRNA close MGC:10272, mRNA, complete cds. /FEA-mRNA close MGC:10272, mRNA, complete cds. /FEA-mRNA laber management laber ma |
| 221918_at | Consensus includes gb:AI742210 /FER-EST /DB_XREF-gi:5110498 /DB_XREF-est:wg39c02.x1 /CLONE-IMAGE:2367458 /UG-HS.183302 E318 |
| 213251_at | 1370 /DB_X |
| 209186_at | gb:M23114.1 /DEF-Homo sapiens calcium-ATPase (HK1) mRNA, complete cds. /FEA-mRNA /ubn-HA1 /Ub_KREF-g1:104100 /UG-h3:1320 Alfase, Carrier transporting, cardiac muscle, slow twitch 2 /FL-gb:M23114.1 |
| 210996_s_at | gb:U43430.1 /DEE-Human epsilon isoform 14-3-3 protein mkNA, Complete cds. /FrAn-manA Fran-11-3-5 protein 15-3-3 protein 15-3 protein 15 |
| 213624_at | Consensus includes gb:AA8/3600 /FEA=E3T /DB_XREF=g1:2959/22 /UB_AREF=8ST:DD4Z4VZ.51 /CLORE-IND6:1020/3 /CC-n3:12340 CC-n3:12340 CC-n3:1234 |
| 204634_at | |
| 212515_s_at | Consensus includes gb:BG492602 /FER=EST /DB XREF=g1:13454114 /DB XREF=eST:00253050661 /CLUNE=IRANGE:105250 /OG-H3:11/10 /LLCH. [Asp-Glu-Ala-AspHis] box polypeptide 3 |
| 218170_at | CGI-111 procein (LOC31013), gb:AF151869.1 gb:NN 016048.1 |
| 210293_s_at | 3 (S. cerevisiae) homolog B, cione Motizago, montr, comptere cus: final montre cus: |
| 221493_at | <pre>gb:AL136629.1 /DEF=Homo sapiens mRNA; cDNA DKF2p564D152 (from clone DKF2p564D152); complete cds. / FEA=mRNA / UEN=UN ED504D152 / PROD=hypothetical protein /DB XREF=gi:12052783 /UG-Hs.278479 / FE-gi:2787-114 / FE-go:2007 / CF-drawn Proteinsse</pre> |
| 213506_at | |
| 201016_at | |
| 203414_at | |

| 213510_x_at | Consensus includes gb:AM194543 /FEA=EST /DB_XREF=gi:6473381 /DB_XREF=est:xb27c02.x1 /CLONE=IMAGE:2577506 /UG-Hs.234573 Homo sapiens mRNA. For |
|-------------|--|
| , | |
| 218539_at | gb:NN 017943.1 /DEF-Homo sapiens hypothetical protein FLJ20725 (FLJ20725), mRNA. /FEA-mRNA /GEM-FLJ20725 /PROD-hypothetical protein |
| | LL020723 / DB XREF=gi:8923650 /UG=Hs.15467 hypothetical protein FLJ20725 /FL=gb:NM 017943.1 |
| 218172_s_at | gb:NM 018630.1 /DEF=Homo sapiens hypothetical protein PRO2577 (PRO2577), mRNR. /FEA-mRNR /GEN-PRO2577 /PROD-hypothetical protein PRO2577 /FL-qb:AF116708.1 qb:NM 018630.1 |
| 202538_s_at | otein (DKF2P564 tein /FL=gb:AF |
| 221504_s_at | |
| | 75 CGI-11 protein /FL=gb:AF298777.1 gb |
| 212213_x_at | Consensus includes gb: AB011139.1 /DEF-Homo sapiens mRNA for KIAA0567 protein, partial cds. /FEA-mRNA /GEN-KIAA0567 /PROD-KIAA0567 protein |
| | 147946 optic atrophy 1 (autosomal dominant) |
| 216652_s_at | Consensus includes gb:AL137673.1 /DEF-Homo sapiens mRNA: cDNA DKF2p434H0872 (from clone DKF2p434H0872). /FEA-mRNA /DB_XREF-g1:6807841 / UG-Hs.306454 Homo sapiens mRNA; cDNA DKF2p434H0672 (from clone DKF2p434H0872) |
| . 221472_at | 3 /DEF-Human DNA sequence from |
| | cerional retinaldehyde-binding protein, the TDE1 gene (Tumour differentially expressed 1), the PKIG gene encoding protein kinase (cAMP-dependent |
| - | 016 /UG=Hs.272168 tumor differentially exp |
| 212526_at | Consensus includes gb:AK002207.1 /DEF=Homo sapiens cDNA FLJ11345 fis, clone PLACE1010877, highly similar to Homo sapiens mRNA for KIAA0610 protein. /FEA=mRNA /DB XREF=qi:7023938 /UG=Hs.118087 KIAA0610 protein. |
| 200624_s_at | Consensus includes gb:AA577695 /FEA=EST /DB_XREF=gi:2355879 /DB_XREF=est:nn22h02.s1 /CLONE~IMAGE:1084659 /UG-Hs.78825 matrin 3 / FI=qb:NM 018834.1 gb:AB018266.1 |
| 201091_s_at | Consensus includes gb: BE748755 /FEA=EST / DB_XREF=g1:10162747 / DB_XREF=est:601571933T1 /CLONE=INAGE:3838737 /UG=HS.278554 heterochromatin- |
| | in 1 /FL-gb:AF136630. |
| 217863_at | Consensus includes gb:AI348378 /FEA=EST /DB_XREF=gi:4085584 /DB_XREF=est:qo20h06.xl /CLONE=IMAGE:1909115 /UG-Hs.75251 DEADH (Asp-Glu-Ala- |
| | nding protein 1 /FL= |
| 202804_at | Consensus includes gb:AI539710 /FEA=EST /DB_XREF=g1:4453845 /DB_XREF=est:tp77b05.x1 /CLONE=IMAGE:2205297 /UG=Hs.89433 ATP-binding casset- |
| | ber 1 /FL-gb: |
| 214198_s_at | Consensus includes gb:AU150824 /FEA=EST /DB_XREF=gi:11012345 /DB_XREF=est:AU150824 /CLONE=NT2RP2003689 /UG=Hs.2491 DiGeorge syndrome |
| • | ene 2 |
| 217976_s_at | gb:NM 016141.1 /DEF=Homo sapiens dynein light chain-A (LOC51143), mRNA. /FEA=mRNA /GEN=LOC51143 /PROD=dynein light chain-A //DB XREP=q1:7705852 / |
| | UG=HS. 266483 dynein 11ght chain-A /FL=gb:AF078849.1 gb:NM 016141.1 |
| 202318_s_at | gb:AF306508.1 /DEF=Homo sapiens SUMO-1 specific protease FKSG6 mRNA, complete cds. /FEA=mRNA /PROD=SUMO-1 specific protease FKSG6 / DE XREF=gi:11096243 /UG-Hs.27197 SUMO-1-specific protease /FL=gb:AF307849.1 gb:AF306508.1 gb:AF196304.1 gb:A |
| 208673_s_at | ıω |
| 218268_at | piens hypothetical protein FLJ12085 (FLJ12085), mRNA. /FEA-mRH |
| | 48827 hypothetical protein FLJ12085 /FL-gb:NM 02277 |
| 209250_at | gb:BC000961.2 /DEF=Homo sapiens, degenerative spermatocyte (homolog Drosophila; lipid desaturase), clone MGC:5079, mRNA, complete cds. / FEA=mRNA /PR00=degenerative spermatocyte (homolog Drosophila; lipid desaturase) /DB_XREF=gi:12803018 /UG=Hs.185973 degenerative spermato- |
| | cyte (homolog Drosophila; lipid desaturase) /EL=qb:BC000961.2 |
| | |

gb:NM 015957.1 /DEF-Homo sapiens CGJ-29 protein (LOCSIO/4), micros / cg-micros / cg-//UG-MS-104058

otein /FL=gb:AF132963.1 gb:NM 015957.1